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# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

# **Patent Application For**

# UNNATURAL REACTIVE AMINO ACID GENETIC CODE ADDITIONS

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# UNNATURAL REACTIVE AMINO ACID GENETIC CODE ADDITIONS

# CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application is a regular utility patent application based on USSN 60/479,931 entitled "Expanding the Eukaryotic Genetic Code" by Chin et al. filed June 18, 2003; USSN 60/493,014 entitled "Expanding the Eukaryotic Genetic Code" by Chin et al., filed August 5, 2003; and USSN 60/496,548 entitled "Expanding the Eukaryotic Genetic Code" by Chin et al., filed August 19, 2003. Priority to and benefit of each of these prior applications is hereby claimed.

# STATEMENT AS TO RIGHTS TO INVENTIONS MADE UNDER FEDERALLY SPONSORED RESEARCH AND DEVELOPMENT

[0002] This invention was made with government support under Grant No. GM 62159 from the National Institutes of Health and support under Grant DE-FG0300ER45812 from the Department of Energy. The government has certain rights to this invention.

#### FIELD OF THE INVENTION

[0003] The invention pertains to the field of translation biochemistry in eukaryotic cells. The invention relates to methods for producing and compositions of orthogonal tRNAs, orthogonal synthetases and pairs thereof, in eukaryotic cells. The invention also relates to compositions of unnatural amino acids, proteins and methods of producing proteins in eukaryotic cells that include unnatural amino acids.

#### **BACKGROUND OF THE INVENTION**

[0004] The genetic code of every known organism, from bacteria to humans, encodes the same twenty common amino acids. Different combinations of the same twenty natural amino acids form proteins that carry out virtually all the complex processes of life, from photosynthesis to signal transduction and the immune response. In order to study and modify protein structure and function, scientists have attempted to manipulate both the genetic code and the amino acid sequence of proteins. However, it has been difficult to remove the constraints imposed by the genetic code that limit proteins to twenty genetically encoded standard building blocks (with the rare exception of selenocysteine (see, e.g., A. Bock et al., (1991), Molecular Microbiology 5:515-20) and pyrrolysine (see, e.g., G. Srinivasan, et al., (2002), Science 296:1459-62).

[0005]Some progress has been made to remove these constraints, although this progress has been limited and the ability to rationally control protein structure and function is still in its infancy. For example, chemists have developed methods and strategies to synthesize and manipulate the structures of small molecules (see, e.g., E. J. Corey, & X.-M. Cheng, The Logic of Chemical Synthesis (Wiley-Interscience, New York, 1995)). Total 5 synthesis (see, e.g., B. Merrifield, (1986), Science 232:341-7 (1986)), and semi-synthetic methodologies (see, e.g., D. Y. Jackson et al., (1994) Science 266:243-7; and, P. E. Dawson, & S. B. Kent, (2000), Annual Review of Biochemistry 69:923-60), have made it possible to synthesize peptides and small proteins, but these methodologies have limited 10 utility with proteins over 10 kilo Daltons (kDa). Mutagenesis methods, though powerful, are restricted to a limited number of structural changes. In a number of cases, it has been possible to competitively incorporate close structural analogues of common amino acids throughout proteins. See, e.g., R. Furter, (1998), Protein Science 7:419-26; K. Kirshenbaum, et al., (2002), ChemBioChem 3:235-7; and, V. Doring et al., (2001), Science 15 292:501-4.

[0006] In an attempt to expand the ability to manipulate protein structure and function, *in vitro* methods using chemically acylated orthogonal tRNAs were developed that allowed unnatural amino acids to be selectively incorporated in response to a nonsense codon, *in vitro* (see, e.g., J. A. Ellman, et al., (1992), Science 255:197-200). Amino acids with novel structures and physical properties were selectively incorporated into proteins to study protein folding and stability and biomolecular recognition and catalysis. See, e.g., D. Mendel, et al., (1995), Annual Review of Biophysics and Biomolecular Structure 24:435-462; and, V. W. Cornish, et al. (Mar. 31, 1995), Angewandte Chemie-International Edition in English 34:621-633. However, the stoichiometric nature of this process severely limited the amount of protein that could be generated.

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[0007] Unnatural amino acids have been microinjected into cells. For example, unnatural amino acids were introduced into the nicotinic acetylcholine receptor in Xenopus oocytes (e.g., M.W. Nowak, et al. (1998), In vivo incorporation of unnatural amino acids into ion channels in Xenopus oocyte expression system, Method Enzymol. 293:504-529) by microinjection of a chemically misacylated Tetrahymena thermophila tRNA (e.g., M.E. Saks, et al. (1996), An engineered Tetrahymena tRNAGln for in vivo incorporation of unnatural amino acids into proteins by nonsense suppression, J. Biol. Chem. 271:23169-23175), and the relevant mRNA. This has allowed detailed biophysical studies of the

receptor in oocytes by the introduction of amino acids containing side chains with unique physical or chemical properties. *See, e.g.*, D.A. Dougherty (2000), *Unnatural amino acids as probes of protein structure and function*, <u>Curr. Opin. Chem. Biol.</u> 4:645-652.

Unfortunately, this methodology is limited to proteins in cells that can be microinjected, and because the relevant tRNA is chemically acylated in vitro, and cannot be re-acylated, the yields of protein are very low.

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[8000]To overcome these limitations, new components were added to the protein biosynthetic machinery of the prokaryote Escherichia coli (E. coli) (e.g., L. Wang, et al., (2001), Science 292:498-500), which allowed genetic encoding of unnatural amino acids in vivo. A number of new amino acids with novel chemical, physical or biological properties, including photoaffinity labels and photoisomerizable amino acids, keto amino acids, and glycosylated amino acids have been incorporated efficiently and with high fidelity into proteins in E. coli in response to the amber codon, TAG, using this methodology. See, e.g., J. W. Chin et al., (2002), Journal of the American Chemical Society 124:9026-9027; J. W. Chin, & P. G. Schultz, (2002), ChemBioChem 11:1135-1137; J. W. Chin, et al., (2002), PNAS United States of America 99:11020-11024: and, L. Wang, & P. G. Schultz, (2002), Chem. Comm., 1-10. However, the translational machinery of prokaryotes and eukaryotes are not highly conserved; thus, components of the biosynthetic machinery added to E.coli cannot often be used to site-specifically incorporate unnatural amino acids into proteins in eukaryotic cells. For example, the Methanococcus jannaschii tyrosyl-tRNA synthetase/tRNA pair that was used in E.coli is not orthogonal in eukaryotic cells. In addition, the transcription of tRNA in eukaryotes, but not in prokaryotes, is carried out by RNA Polymerase III and this places restrictions on the primary sequence of the tRNA structural genes that can be transcribed in eukaryotic cells. Moreover, in contrast to prokaryotic cells, tRNAs in eukaryotic cells need to be exported from the nucleus, where they are transcribed, to the cytoplasm, to function in translation. Finally, the eukaryotic 80S ribosome is distinct from the 70S prokaryotic ribosome. Thus, there is a need to develop improved components of the biosynthetic machinery to expand the eukaryotic genetic code. This invention fulfills these and other needs, as will be apparent upon review of the following disclosure.

#### SUMMARY OF THE INVENTION

[0009] The invention provides eukaryotic cells with translation components, e.g., pairs of orthogonal aminoacyl-tRNA synthetases (O-RSs) and orthogonal tRNAs (O-tRNAs) and individual components thereof, that are used in eukaryotic protein biosynthetic machinery to incorporate an unnatural amino acid in a growing polypeptide chain, in a eukaryotic cell.

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[0010] Compositions of the invention include a eukaryotic cell (e.g., a yeast cell (such as a *Saccharomyces cerevisiae* cell), a mammalian cell, a plant cell, an algae cell, a fungal cell, an insect cell, etc.) comprising an orthogonal aminoacyl-tRNA synthetase (O-RS) (e.g., derived from a non-eukaryotic organism, such as *Escherichia coli, Bacillus stearothermophilus*, etc.), where the O-RS preferentially aminoacylates an orthogonal tRNA (O-tRNA) with at least one unnatural amino acid in the eukaryotic cell. Optionally, two or more OtRNAs can be aminoacylated in a given eukaryotic cell. In one aspect, an O-RS aminoacylates an O-tRNA with the unnatural amino acid, e.g., at least 40%, at least 45%, at least 50%, at least 60%, at least 75%, at least 80%, or even 90% or more as efficiently as does an O-RS having an amino acid sequence, e.g., as set forth in SEQ ID NO.: 86 or 45. In one embodiment, an O-RS of the invention aminoacylates the O-tRNA with the unnatural amino acid, e.g., at least 10-fold, at least 20-fold, at least 30-fold, etc., more efficiently than the O-RS aminoacylates the O-tRNA with a natural amino acid.

[0011] In one embodiment, the O-RS or a portion thereof is encoded by a polynucleotide sequence as set forth in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), or a complementary polynucleotide sequence thereof. In another embodiment, the O-RS comprises an amino acid sequence as set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, or a conservative variation thereof. In yet another embodiment, the O-RS comprises an amino acid sequence that is, e.g., at least 90%, at least 95%, at least 98%, at least 99%, or at least 99.5 % or more, identical to that of a naturally occurring tyrosyl aminoacyl-tRNA synthetase (TyrRS) and comprises two or more amino acids from groups A-E. Group A includes valine, isoleucine, leucine, glycine, serine, alanine, or threonine at a position corresponding to Tyr37 of an *E. coli* TyrRS. Group B includes aspartate at a position corresponding to Asn126 of an *E. coli* TyrRS. Group C includes threonine, serine, arginine, asparagine or glycine at a position corresponding to Asp182 of an *E. coli* TyrRS. Group D includes methionine, alanine, valine, or tyrosine at a position corresponding to Phe183 of an

E. coli TyrRS; and, group E includes serine, methionine, valine, cysteine, threonine, or alanine at a position corresponding to Leu186 of an E. coli TyrRS.

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[0012] Any subset of combinations of these groups are a feature of the invention. For example, in one embodiment, the O-RS has two or more amino acids selected from valine, isoleucine, leucine, or threonine occurs at a position corresponding to Tyr37 of *E. coli* TyrRS; threonine, serine, arginine, or glycine at a position corresponding to Asp182 of *E. coli* TyrRS; methionine, or tyrosine at a position corresponding to Phe183 of *E. coli* TyrRS. In another embodiment, the O-RS includes two more more amino acids selected from glycine, serine, or alanine at a position corresponding to Tyr37 of *E. coli* TyrRS, aspartate at a position corresponding to Asp182 of *E. coli* TyrRS, alanine, or valine, at a position corresponding to Phe183 of *E. coli* TyrRS, and/or methionine, valine, cysteine, or threonine, at a position corresponding to Leu186 of *E. coli* TyrRS.

15 [0013] In another embodiment, the O-RS has one or more improved or enhanced enzymatic properties for the unnatural amino acid as compared to a natural amino acid. For example, the improved or enhanced properties for the unnatural amino acid as compared to a natural amino acid include any of, e.g., a higher Km, a lower Km, a higher kcat, a lower kcat, a lower kcat/km, a higher kcat/km, etc.

[0014] The eukaryotic cell also optionally includes an unnatural amino acid(s). The eukaryotic cell optionally includes an orthogonal tRNA (O-tRNA) (e.g., derived from a non-eukaryotic organism, such as *Escherichia coli*, *Bacillus stearothermophilus*, and/or the like), where the O-tRNA recognizes a selector codon and is preferentially aminoacylated with the unnatural amino acid by the O-RS. In one aspect, the O-tRNA mediates the incorporation of the unnatural amino acid into a protein with, e.g., at least 45%, at least 50%, at least 60%, at least 75%, at least 80%, at least 90%, at least 95%, or 99% or the efficiency of a tRNA that comprises or is processed in a cell from a polynucleotide sequence as set forth in SEQ ID NO.: 65. In another aspect, the O-tRNA comprises the sequence of SEQ ID NO.:65, and the O-RS comprises a polypeptide sequence selected from an amino acid sequence set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, and/or a conservative variation thereof.

[0015] In another embodiment, the eukaryotic cell comprises a nucleic acid that comprises a polynucleotide that encodes a polypeptide of interest, where the polynucleotide comprises a selector codon that is recognized by the O-tRNA. In one aspect, the yield of the polypeptide of interest comprising the unnatural amino acid is, e.g., at least 2.5%, at least 5%, at least 10%, at least 25%, at least 30%, at least 40%, 50% or more, of that obtained for the naturally occurring polypeptide of interest from a cell in which the polynucleotide lacks the selector codon. In another aspect, the cell produces the polypeptide of interest in the absence of the unnatural amino acid, with a yield that is, e.g., less than 35%, less than 30%, less than 20%, less than 15%, less than 10%, less than 5%, less than 2.5%, etc., of the yield of the polypeptide in the presence of the unnatural amino acid.

[0016] The invention also provides a eukaryotic cell comprising an orthogonal aminoacyl-tRNA synthetase (O-RS), an orthogonal tRNA (O-tRNA), an unnatural amino acid, and a nucleic acid that comprises a polynucleotide that encodes a polypeptide of interest. The polynucleotide comprises a selector codon that is recognized by the O-tRNA. In addition, the O-RS preferentially aminoacylates the orthogonal tRNA (O-tRNA) with the unnatural amino acid in the eukaryotic cell, and the cell produces the polypeptide of interest in the absence of the unnatural amino acid, with a yield that is, e.g., less than 30%, less than 20%, less than 15%, less than 5%, less than 2.5%, etc., of the yield of the polypeptide in the presence of the unnatural amino acid.

[0017] Compositions that include a eukaryotic cell comprising an orthogonal tRNA (O-tRNA) are also a feature of the invention. Typically, the O-tRNA mediates incorporation of an unnatural amino acid into a protein that is encoded by a polynucleotide that comprises a selection codon that is recognized by the O-tRNA in vivo. In one embodiment, the O-tRNA mediates the incorporation of the unnatural amino acid into the protein with, e.g., at least 45%, at least 50%, at least 60%, at least 75%, at least 80%, at least 90%, at least 95%, or even 99% or more the efficiency of a tRNA that comprises or is processed in a cell from a polynucleotide sequence as set forth in SEQ ID NO.: 65. In another embodiment, the O-tRNA comprises or is processed from a polynucleotide sequence as set forth in SEQ ID NO.: 65, or a conservative variation thereof. In yet another embodiment, the O-tRNA comprises a recyclable O-tRNA.

[0018] In one aspect of the invention, the O-tRNA is post-transcriptionally modified. The invention also provides a nucleic acid that encodes an O-tRNA in a eukaryotic cell, or a complementary polynucleotide thereof. In one embodiment, the nucleic acid comprises an A box and a B box.

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The invention also features methods of producing translational components, [0019] e.g., O-RSs or O-tRNA/O-RS pairs (and translational components produced by these methods). For example, the invention provides methods of producing an orthogonal aminoacyl-tRNA synthetase (O-RS) that preferentially aminoacylates an orthogonal tRNA with an unnatural amino acid in a eukaryotic cell. The method includes, e.g., (a) subjecting to positive selection, in the presence of an unnatural amino acid, a population of eukaryotic cells of a first species, where the eukaryotic cells each comprise: i) a member of a library of aminoacyl-tRNA synthetases (RSs), ii) an orthogonal tRNA (O-tRNA), iii) a polynucleotide that encodes a positive selection marker, and iv) a polynucleotide that encodes a negative selection marker; where cells that survive the positive selection comprise an active RS that aminoacylates the orthogonal tRNA (O-tRNA) in the presence of an unnatural amino acid. The cells that survive the positive selection are subjected to negative selection in the absence of the unnatural amino acid to eliminate active RSs that aminoacylate the O-tRNA with a natural amino acid. This provides the O-RS that preferentially aminoacylates the OtRNA with the unnatural amino acid.

[0020] In certain embodiments, the polynucleotide that encodes the positive selection marker is operably linked to a response element and the cells further comprise a polynucleotide that: a) encodes a transcriptional modulator protein (e.g., a eukaryotic transcriptional modulator protein, etc.) that modulates transcription from the response element, and b) comprises at least one selector codon. The incorporation of the unnatural amino acid into the transcriptional modulator protein by the O-tRNA aminoacylated with the unnatural amino acid results in transcription of the positive selection marker. In one embodiment, the transcriptional modulator protein is a transcriptional activator protein (e.g., GAL4, etc.), and the selector codon is an amber stop codon, e.g., where the amber stop codon is located in or substantially near a portion of the polynucleotide that encodes a DNA binding domain of the transcriptional activator protein.

[0021] The positive selection marker can be any of a variety of molecules. In one embodiment, the positive selection marker comprises a nutritional supplement for growth

and the selection is performed on a medium that lacks the nutritional supplement. In another embodiment, the polynucleotide that encodes the positive selection marker is, e.g., an ura3, leu2, lys2, lacZ gene, his3 (e.g., where the his3 gene encodes an imidazole glycerol phosphate dehydratase, detected by providing 3-aminotriazole (3-AT)), and/or the like. In yet another embodiment, the polynucleotide that encodes the positive selection marker comprises a selector codon.

[0022] As with the positive selection marker, the negative selection marker can also be any of a variety of molecules. In certain embodiments, the polynucleotide that encodes the negative selection marker is operably linked to a response element from which transcription is mediated by the transcriptional modulator protein. The incorporation of a natural amino acid into the transcriptional modulator protein by the O-tRNA aminoacylated with a natural amino acid results in transcription of the negative selection marker. In one embodiment, the polynucleotide that encodes the negative selection marker is, e.g., an ura3 gene and the negative selection is accomplished on a medium that comprises 5-fluroorotic acid (5-FOA). In another embodiment, the medium used for negative selection comprises a selecting or screening agent that is converted to a detectable substance by the negative selection marker. In one aspect of the invention, the detectable substance is a toxic substance. In one embodiment, the polynucleotide that encodes the negative selection marker comprises a selector codon.

[0023] In certain embodiments, the positive selection marker and/or the negative selection marker comprises a polypeptide that fluoresces or catalyzes a luminescent reaction in the presence of a suitable reactant. In one aspect of the invention, the positive selection marker and/or the negative selection marker is detected by fluorescence-activated cell sorting (FACS), or by luminescence. In certain embodiments, the positive selection marker and/or negative selection marker comprises an affinity based screening marker, or a transcriptional modulator protein. In one embodiment, the same polynucleotide encodes both the positive selection marker and the negative selection marker.

[0024] In one embodiment, the polynucleotide that encodes the positive selection marker and/or negative selection marker of the invention can comprises at least two selector codons, which each or both can comprise at least two different selector codons or at least two of the same selector codons.

[0025] Additional levels of selection/screening stringency can also be used in the methods of the invention. In one embodiment, the methods can comprise, e.g., providing a varying amount of an inactive synthetase in step (a), (b) or both (a) and (b), where the varying amount of the inactive synthetase provides an additional level of selection or screening stringency. In one embodiment, step (a), (b) or both steps (a) and (b) of the method for producing an O-RS includes varying a selection or screening stringency, e.g., of the positive and/or negative selection marker. The method optionally includes subjecting the O-RS that preferentially aminoacylates the O-tRNA with the unnatural amino acid to an additional selection round, e.g., an additional positive selection round(s), an additional negative selection round(s) or combinations of both additional positive and negative selection rounds.

[0026] In one embodiment, the selecting/screening comprises one or more positive or negative selection/screening chosen from, e.g., a change in amino acid permeability, a change in translation efficiency, a change in translational fidelity, etc. The one or more change is based upon a mutation in one or more polynucleotide that encodes a component of orthogonal tRNA-tRNA synthetase pair is used to produce protein.

[0027] Typically, the library of RSs (e.g., a library of mutant RSs) comprises RSs derived from at least one aminoacyl-tRNA synthetase (RS), e.g., from a non-eukaryotic organism. In one embodiment, the library of RSs is derived from an inactive RS, e.g., where the inactive RS is generated by mutating an active RS. In another embodiment, the inactive RS comprises an amino acid binding pocket and one or more amino acids that comprise the binding pocket are substituted with one or more different amino acids, e.g., the substituted amino acids are substituted with alanines.

[0028] In certain embodiments, the method of producing an O-RS further includes performing random mutation, site-specific mutation, recombination, chimeric construction, or any combination thereof, on a nucleic acid that encodes an RS, thereby producing the library of mutant RSs. In certain embodiments, the method further includes, e.g., (c) isolating a nucleic acid that encodes the O-RS; (d) generating from the nucleic acid a set of polynucleotides that encode mutated O-RSs (e.g., by random mutagenesis, site-specific mutagenesis, chimeric construction, recombination or any combination thereof); and, (e) repeating steps (a) and/or (b) until a mutated O-RS is obtained that preferentially

aminoacylates the O-tRNA with the unnatural amino acid. In one aspect of the invention, steps (c)-(e) are performed at least two times.

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[0029] Methods of producing O-tRNA/O-RS pairs are also a feature of the invention. In one embodiment, the O-RS is obtained as described above and the O-tRNA is obtained by subjecting to negative selection a population of eukaryotic cells of a first species, where the eukaryotic cells comprise a member of a library of tRNAs, to eliminate cells that comprise a member of the library of tRNAs that is aminoacylated by an aminoacyl-tRNA synthetase (RS) that is endogenous to the eukaryotic cells. This provides a pool of tRNAs that are orthogonal to the eukaryotic cell of the first species. In one aspect of the invention, the library of tRNAs comprises tRNAs derived from at least one tRNA, e.g., from a non-eukaryotic organism. In another aspect of the invention, the library of aminoacyl-tRNA synthetases (RSs) comprises RSs derived from at least one aminoacyltRNA synthetase (RS), e.g., from a non-eukaryotic organism. In yet another aspect of the invention, the library of tRNAs comprises tRNAs derived from at least one tRNA from a first non-eukaryotic organism. The library of aminoacyl-tRNA synthetases (RSs) optionally comprises RSs derived from at least one aminoacyl-tRNA synthetase (RS) from a second non-eukaryotic organism. In one embodiment, the first and second non-eukaryotic organisms are the same. Alternatively, the first and second non-eukaryotic organisms can be different. Specific O-tRNA/O-RS pairs produced by the methods of the invention are also a feature of the invention.

[0030] Another feature of the invention is a method for producing translational components in one species and introducing the selected/screened translational components into a second species. For example, the method of producing a O-tRNA/O-RS pair in a first species (e.g., a eukaryotic species, such as a yeast and the like) further includes introducing a nucleic acid that encodes the O-tRNA and a nucleic acid that encodes the O-RS into a eukaryotic cell of a second species (e.g., a mammal, an insect, a fungus, an algae, a plant and the like). The second species can use the introduced translational components to incorporate an unnatural amino acid into a growing polypeptide chain in vivo, e.g., during translation.

[0031] In another example, a method of producing an orthogonal aminoacyl-tRNA synthetase (O-RS) that preferentially aminoacylates an orthogonal tRNA with an unnatural amino acid in a eukaryotic cell includes: (a) subjecting to positive selection, in the presence

of an unnatural amino acid, a population of eukaryotic cells of a first species (e.g., a eukaryotic species, such as a yeast or the like). The eukaryotic cells of the first species each comprise: i) a member of a library of aminoacyl-tRNA synthetases (RSs), ii) an orthogonal tRNA (O-tRNA), iii) a polynucleotide that encodes a positive selection marker, and iv) a polynucleotide that encodes a negative selection marker. The cells that survive the positive selection comprise an active RS that aminoacylates the orthogonal tRNA (O-tRNA) in the presence of an unnatural amino acid. The cells that survive the positive selection are subjected to negative selection in the absence of the unnatural amino acid to eliminate active RSs that aminoacylate the O-tRNA with a natural amino acid, thereby providing an O-RS that preferentially aminoacylates the O-tRNA with the unnatural amino acid. A nucleic acid that encodes the O-tRNA and a nucleic acid that encodes the O-RS are introduced into a eukaryotic cell of a second species (e.g., mammal, an insect, a fungus, an algae, a plant and/or the like). These components, when translated in the second species, can be used to incorporate unnatural amino acids into a protein or polypeptide of interest in the second species. In one embodiment, the O-tRNA and/or the O-RS are introduced into a eukaryotic cell of a second species.

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[0032] In certain embodiments, the O-tRNA is obtained by subjecting to negative selection a population of eukaryotic cells of a first species, where the eukaryotic cells comprise a member of a library of tRNAs, to eliminate cells that comprise a member of the library of tRNAs that is aminoacylated by an aminoacyl-tRNA synthetase (RS) that is endogenous to the eukaryotic cells. This provides a pool of tRNAs that are orthogonal to the eukaryotic cell of the first species and the second species.

[0033] In one aspect, the invention comprises a composition comprising a protein, wherein the protein comprises at least one unnatural amino acid and at least one post-translational modification, wherein the at least one post-translational modification comprises attachment of a molecule comprising a second reactive group by a [3+2] cycloaddition to the at least one unnatural amino acid comprising a first reactive group.

[0034] Thus, proteins (or polypeptides of interest) with at least one unnatural amino acid are also a feature of the invention. In certain embodiments of the invention, a protein with at least one unnatural amino acid includes at least one post-translational modification. In one embodiment, the at least one post-translational modification comprises attachment of a molecule (e.g., a dye, a polymer, e.g., a derivative of polyethylene glycol, a

photocrosslinker, a cytotoxic compound, an affinity label, a derivative of biotin, a resin, a second protein or polypeptide, a metal chelator, a cofactor, a fatty acid, a carbohydrate, a polynucleotide (e.g., DNA, RNA, etc.), etc.) comprising a second reactive group by a [3+2] cycloaddition to the at least one unnatural amino acid comprising a first reactive group. For example, the first reactive group is an alkynyl moiety (e.g., in the unnatural amino acid *p*-propargyloxyphenylalanine) (this group is also sometimes refer to as an acetylene moiety) and the second reactive group is an azido moiety. In another example, the first reactive group is the azido moiety (e.g., in the unnatural amino acid *p*-azido-L-phenylalanine) and the second reactive group is the alkynyl moiety. In certain embodiments, a protein of the invention includes at least one unnatural amino acid (e.g., a keto unnatural amino acid) comprising at least one post-translational modification, where the at least one post-translational modification comprises a saccharide moiety. In certain embodiments, the post-translational modification is made in vivo in a eukaryotic cell.

[0035] In certain embodiments, the protein includes at least one post-translational modification that is made in vivo by a eukaryotic cell, where the post-translational modification is not made by a prokaryotic cell. Examples of post-translational modifications include, but are not limited to, acetylation, acylation, lipid-modification, palmitoylation, palmitate addition, phosphorylation, glycolipid-linkage modification, and the like. In one embodiment, the post-translational modification comprises attachment of an oligosaccharide to an asparagine by a GlcNAc-asparagine linkage (e.g., where the oligosaccharide comprises (GlcNAc-Man)<sub>2</sub>-Man-GlcNAc-GlcNAc, and the like). In another embodiment, the post-translational modification comprises attachment of an oligosaccharide (e.g., Gal-GalNAc, Gal-GlcNAc, etc.) to a serine or threonine by a GalNAc-serine, a GalNAc-threonine, a GlcNAc-serine, or a GlcNAc-threonine linkage. In certain embodiments, a protein or polypeptide of the invention can comprise a secretion or localization sequence, an epitope tag, a FLAG tag, a polyhistidine tag, a GST fusion, and/or the like.

[0036] Typically, the proteins are, e.g., at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, at least 95%, or even at least 99% or more identical to any available protein (e.g., a therapeutic protein, a diagnostic protein, an industrial enzyme, or portion thereof, and/or the like), and they comprise one or more unnatural amino acid. In one embodiment, a composition of the invention includes a protein or polypeptide of interest and an excipient (e.g., a buffer, a pharmaceutically acceptable excipient, etc.).

[0037] The protein or polypeptide of interest can contain at least one, at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, or ten or more unnatural amino acids. The unnatural amino acids can be the same or different, e.g., there can be 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or more different sites in the protein that comprise 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or more different unnatural amino acids. In certain embodiments, at least one, but fewer than all, of a particular amino acid present in a naturally occurring version of the protein is substituted with an unnatural amino acid.

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Examples of a protein (or polypeptide of interest) include, but are not limited [0038]to, e.g., a cytokine, a growth factor, a growth factor receptor, an interferon, an interleukin, an inflammatory molecule, an oncogene product, a peptide hormone, a signal transduction molecule, a steroid hormone receptor, erythropoietin (EPO), insulin, human growth hormone, an Alpha-1 antitrypsin, an Angiostatin, an Antihemolytic factor, an antibody, an Apolipoprotein, an Apoprotein, an Atrial natriuretic factor, an Atrial natriuretic polypeptide, an Atrial peptide, a C-X-C chemokine, T39765, NAP-2, ENA-78, a Gro-a, a Gro-b, a Gro-c, an IP-10, a GCP-2, an NAP-4, an SDF-1, a PF4, a MIG, a Calcitonin, a c-kit ligand, a cytokine, a CC chemokine, a Monocyte chemoattractant protein-1, a Monocyte chemoattractant protein-2, a Monocyte chemoattractant protein-3, a Monocyte inflammatory protein-1 alpha, a Monocyte inflammatory protein-1 beta, RANTES, I309, R83915, R91733, HCC1, T58847, D31065, T64262, a CD40, a CD40 ligand, a C-kit Ligand, a Collagen, a Colony stimulating factor (CSF), a Complement factor 5a, a Complement inhibitor, a Complement receptor 1, a cytokine, DHFR, an epithelial Neutrophil Activating Peptide-78, a GROα/MGSA, a GROβ, a GROγ a MIP-1α, a MIP-1δ, a MCP-1, an Epidermal Growth Factor (EGF), an epithelial Neutrophil Activating Peptide, an Erythropoietin (EPO), an Exfoliating toxin, a Factor IX, a Factor VII, a Factor VIII, a Factor X, a Fibroblast Growth Factor (FGF), a Fibringen, a Fibronectin, a G-CSF, a GM-CSF, a Glucocerebrosidase, a Gonadotropin, a growth factor, a growth factor receptor, a Hedgehog protein, a Hemoglobin, a Hepatocyte Growth Factor (HGF), a Hirudin, a Human serum albumin, an ICAM-1, an ICAM-1 receptor, an LFA-1, an LFA-1 receptor, an Insulin, an Insulin-like Growth Factor (IGF), an IGF-I, an IGF-II, an interferon, an IFN-α, an IFNβ, an IFN-γ, an interleukin, an IL-1, an IL-2, an IL-3, an IL-4, an IL-5, an IL-6, an IL-7, an IL-8, an IL-9, an IL-10, an IL-11, an IL-12, a Keratinocyte Growth Factor (KGF), a Lactoferrin, a leukemia inhibitory factor, a Luciferase, a Neurturin, a Neutrophil inhibitory factor (NIF), an oncostatin M, an Osteogenic protein, an oncogene product, a Parathyroid

hormone, a PD-ECSF, a PDGF, a peptide hormone, a Human Growth Hormone, a Pleiotropin, a Protein A, a Protein G, a Pyrogenic exotoxins A, B, or C, a Relaxin, a Renin, an SCF, a Soluble complement receptor I, a Soluble I-CAM 1, a Soluble interleukin receptors, a Soluble TNF receptor, a Somatomedin, a Somatostatin, a Somatotropin, a 5 Streptokinase, a Superantigens, a Staphylococcal enterotoxins, an SEA, an SEB, an SEC1, an SEC2, an SEC3, an SED, an SEE, a steroid hormone receptor, a Superoxide dismutase (SOD), a Toxic shock syndrome toxin, a Thymosin alpha 1, a Tissue plasminogen activator, a tumor growth factor (TGF), a TGF-α, a TGF-β, a Tumor Necrosis Factor, a Tumor Necrosis Factor alpha, a Tumor necrosis factor beta, a Tumor necrosis factor receptor 10 (TNFR), a VLA-4 protein, a VCAM-1 protein, a Vascular Endothelial Growth Factor (VEGEF), a Urokinase, a Mos, a Ras, a Raf, a Met; a p53, a Tat, a Fos, a Myc, a Jun, a Myb, a Rel, an estrogen receptor, a progesterone receptor, a testosterone receptor, an aldosterone receptor, an LDL receptor, a SCF/c-Kit, a CD40L/CD40, a VLA-4/VCAM-1, an ICAM-1/LFA-1, a hyalurin/CD44, a corticosterone, a protein present in Genebank or 15 other available databases, and the like, and/or a portion thereof. In one embodiment, the polypeptide of interest includes a transcriptional modulator protein (e.g., a transcriptional activator protein (such as GAL4), or a transcriptional repressor protein, etc.) or a portion thereof.

[0039] Compositions of a GAL4 protein, or portion thereof, in a eukaryotic cell are also a feature of the invention. Typically, the GAL4 protein or portion thereof comprises at least one unnatural amino acid.

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[0040] A eukaryotic cell of the invention provides the ability to synthesize proteins that comprise unnatural amino acids in large useful quantities. For example, proteins comprising an unnatural amino acid can be produced at a concentration of, e.g., at least 10 μg/liter, at least 50 μg/liter, at least 75 μg/liter, at least 100 μg/liter, at least 200 μg/liter, at least 250 μg/liter, or at least 500 μg/liter or more of protein in a cell extract, a buffer, a pharmaceutically acceptable excipient, and/or the like. In certain embodiments, a composition of the invention includes, e.g., at least 10 μg, at least 50μg, at least 75 μg, at least 100 μg, at least 200 μg, at least 250 μg, or at least 500μg or more of protein that comprises a unnatural amino acid.

[0041] In certain embodiments, the protein or polypeptide of interest (or portion thereof) is encoded by a nucleic acid. Typically, the nucleic acid comprises at least one

selector codon, at least two selector codons, at least three selector codons, at least four selector codons, at least five selector codons, at least six selector codons, at least seven selector codons, at least eight selector codons, at least nine selector codons, or even ten or more selector codons.

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[0042] The invention also provides methods for producing, in a eukaryotic cell, at least one protein comprising at least one unnatural amino acid (as well as proteins produced by such methods). The methods include, e.g., growing, in an appropriate medium, a eukaryotic cell that comprises a nucleic acid that comprises at least one selector codon and encodes the protein. The eukaryotic cell also comprises an orthogonal tRNA (O-tRNA) that functions in the cell and recognizes the selector codon and an orthogonal aminoacyl tRNA synthetase (O-RS) that preferentially aminoacylates the O-tRNA with the unnatural amino acid, and the medium comprises an unnatural amino acid. In one embodiment, the O-RS aminoacylates the O-tRNA with the unnatural amino acid e.g., at least 45%, at least 50%, at least 60%, at least 75%, at least 80%, at least 90%, at least 95%, or even 99% or more as efficiently as does an O-RS having an amino acid sequence, e.g., as set forth in SEQ ID NO.: 86 or 45. In another embodiment, the O-tRNA comprises, is processed from, or is encoded by SEQ ID NO.: 64 or 65, or a complementary polynucleotide sequence thereof. In yet another embodiment, the O-RS comprises an amino acid sequence as set forth in any one of SEO ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86.

In one embodiment, the method further includes incorporating into the protein the unnatural amino acid, where the unnatural amino acid comprises a first reactive group; and contacting the protein with a molecule (e.g., a dye, a polymer, e.g., a derivative of polyethylene glycol, a photocrosslinker, a cytotoxic compound, an affinity label, a derivative of biotin, a resin, a second protein or polypeptide, a metal chelator, a cofactor, a fatty acid, a carbohydrate, a polynucleotide (e.g., DNA, RNA, etc.), etc.) that comprises a second reactive group. The first reactive group reacts with the second reactive group to attach the molecule to the unnatural amino acid through a [3+2] cycloaddition. In one embodiment, the first reactive group is an alkynyl or azido moiety and the second reactive group is an azido or alkynyl moiety. For example, the first reactive group is the alkynyl moiety (e.g., in unnatural amino acid p-propargyloxyphenylalanine) and the second reactive group is the azido moiety. In another example, the first reactive group is the azido moiety (e.g., in the unnatural amino acid p-azido-L-phenylalanine) and the second reactive group is the alkynyl moiety.

In certain embodiments, the encoded protein comprises a therapeutic protein, a diagnostic protein, an industrial enzyme, or portion thereof. In one embodiment, the protein that is produced by the method is further modified through the unnatural amino acid. For example, the unnatural amino acid is modified through, e.g., a nucleophilic-electrophilic reaction, through a [3+2] cycloaddition, etc. In another embodiment, the protein produced by the method is modified by at least one post-translational modification (e.g., N-glycosylation, O-glycosylation, acetylation, acylation, lipid-modification, palmitoylation, palmitate addition, phosphorylation, glycolipid-linkage modification, and the like) in vivo.

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Methods of producing a screening or selecting transcriptional modulator [0045] protein are also provided (as are screening or selecting transcriptional modulator proteins produced by such methods). The methods include, e.g., selecting a first polynucleotide sequence, where the polynucleotide sequence encodes a nucleic acid binding domain; and mutating the first polynucleotide sequence to include at least one selector codon. This provides a screening or selecting polynucleotide sequence. The methods also include, e.g., selecting a second polynucleotide sequence, where the second polynucleotide sequence encodes a transcriptional activation domain; providing a construct that comprises the screening or selecting polynucleotide sequence operably linked to the second polynucleotide sequence; and, introducing the construct, an unnatural amino acid, an orthogonal tRNA synthetase (O-RS) and an orthogonal tRNA (O-tRNA), into a cell. With these components, the O-RS preferentially aminoacylates the O-tRNA with the unnatural amino acid and the O-tRNA recognizes the selector codon and incorporates the unnatural amino acid into the nucleic acid binding domain, in response to the selector codon in the screening or selecting polynucleotide sequence. This provides the screening or selecting transcriptional modulator protein.

[0046] In certain embodiments, the compositions and the methods of the invention include eukaryotic cells. A eukaryotic cell of the invention includes any of, e.g., a mammalian cell, a yeast cell, a fungus cell, a plant cell, an insect cell, etc. The translation components of the invention can be derived from a variety of organisms, e.g., non-eukaryotic organisms, such as a prokaryotic organism (e.g., E. coli, Bacillus stearothermophilus, or the like), or an archaebacterium, or e.g., a eukaryotic organism.

[0047] A selector codon of the invention expands the genetic codon framework of eukaryotic protein biosynthetic machinery. Any of a variety of selector codons can be used

in the invention, including stop codons (e.g., an amber codon, an ochre codon, or an opal stop codon), nonsense codons, rare codons, four (or more) base codons, and/or the like.

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[0048] Examples of unnatural amino acids that can be used in the compositions and methods described herein include (but are not limited to): a p-acetyl-L-phenylalanine, a piodo-L-phenylalanine, an O-methyl-L-tyrosine, a p-propargyloxyphenylalanine, a ppropargyl-phenylalanine, an L-3-(2-naphthyl)alanine, a 3-methyl-phenylalanine, an O-4allyl-L-tyrosine, a 4-propyl-L-tyrosine, a tri-O-acetyl-GlcNAcβ-serine, an L-Dopa, a fluorinated phenylalanine, an isopropyl-L-phenylalanine, a p-azido-L-phenylalanine, a pacyl-L-phenylalanine, a p-benzoyl-L-phenylalanine, an L-phosphoserine, a phosphonoserine, a phosphonotyrosine, a p-bromophenylalanine, a p-amino-L-phenylalanine, an isopropyl-Lphenylalanine, an unnatural analogue of a tyrosine amino acid; an unnatural analogue of a glutamine amino acid; an unnatural analogue of a phenylalanine amino acid; an unnatural analogue of a serine amino acid; an unnatural analogue of a threonine amino acid; an alkyl, aryl, acyl, azido, cyano, halo, hydrazine, hydrazide, hydroxyl, alkenyl, alkynl, ether, thiol, sulfonyl, seleno, ester, thioacid, borate, boronate, phospho, phosphono, phosphine, heterocyclic, enone, imine, aldehyde, hydroxylamine, keto, or amino substituted amino acid, or any combination thereof; an amino acid with a photoactivatable cross-linker; a spinlabeled amino acid; a fluorescent amino acid; a metal binding amino acid; a metalcontaining amino acid; a radioactive amino acid; a photocaged and/or photoisomerizable amino acid; a biotin or biotin-analogue containing amino acid; a keto containing amino acid; an amino acid comprising polyethylene glycol or polyether; a heavy atom substituted amino acid; a chemically cleavable or photocleavable amino acid; an amino acid with an elongated side chain; an amino acid containing a toxic group; a sugar substituted amino acid; a carbon-linked sugar-containing amino acid; a redox-active amino acid; an α-hydroxy containing acid; an amino thio acid; an  $\alpha$ ,  $\alpha$  disubstituted amino acid; a  $\beta$ -amino acid; a cyclic amino acid other than proline or histidine, an aromatic amino acid other than phenylalanine, tyrosine or tryptophan, and/or the like.

[0049] The invention also provides polypeptides (O-RSs) and polynucleotides, e.g., O-tRNAs, polynucleotides that encode O-RSs or portions thereof (e.g., the active site of the synthetase), oligonucleotides used to construct aminoacyl-tRNA synthetase mutants, polynucleotides that encode a protein or polypeptide of interest that comprise one or more selector codon, etc. For example, a polypeptide of the invention includes a polypeptide that

comprises an amino acid sequence as set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, a polypeptide that comprises an amino acid sequence encoded by a polynucleotide sequence as set forth in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), and a polypeptide that is specifically immunoreactive with an antibody specific for a polypeptide that comprises an amino acid sequence as shown in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, or a polypeptide that comprises an amino acid sequence encoded by a polynucleotide sequence as shown in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35).

[0050] Also included among the polypeptides of the invention is a polypeptide that comprises an amino acid sequence that is at least 90% identical to that of a naturally occurring tyrosyl aminoacyl-tRNA synthetase (TyrRS) (e.g., SEQ ID NO.:2) and comprises two or more amino acids of groups A-E (noted above). Similarly, polypeptides of the invention also optionally include a polypeptide that comprises at least 20 contiguous amino acids of any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, and two or more amino acid substitutions as indicated above in groups A-E. An amino acid sequence comprising a conservative variation of any of the above polypeptides is also included as a polypeptide of the invention.

[0051] In one embodiment, a composition includes a polypeptide of the invention and an excipient (e.g., buffer, water, pharmaceutically acceptable excipient, etc.). The invention also provides an antibody or antisera specifically immunoreactive with a polypeptide of the invention.

[0052] Polynucleotides are also provided in the invention. Polynucleotides of the invention include those that encode proteins or polypeptides of interests of the invention with one or more selector codon. In addition, polynucleotides of the invention include, e.g., a polynucleotide comprising a nucleotide sequence as set forth in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), 64-85; a polynucleotide that is complementary to or that encodes a polynucleotide sequence thereof; and/or a polynucleotide encoding a polypeptide that comprises an amino acid sequence as set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, or a conservative variation thereof. A polynucleotide of the invention also includes a polynucleotide that encodes a polypeptide of the invention. Similarly, a nucleic acid that

hybridizes to a polynucleotide indicated above under highly stringent conditions over substantially the entire length of the nucleic acid is a polynucleotide of the invention.

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[0053] A polynucleotide of the invention also includes a polynucleotide that encodes a polypeptide that comprises an amino acid sequence that is at least 90% identical to that of a naturally occurring tyrosyl aminoacyl-tRNA synthetase (TyrRS) (e.g., SEQ ID NO.: 2) and comprises two or more mutations as indicated above in groups A-E (noted above). A polynucleotide that is that is at least 70%, (or at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 98%, or least 99% or more) identical to a polynucleotide indicated above and/or a polynucleotide comprising a conservative variation of any of the polynucleotides indicated above are also included among the polynucleotides of the invention.

[0054] In certain embodiments, a vector (e.g., a plasmid, a cosmid, a phage, a virus, etc.) comprises a polynucleotide of the invention. In one embodiment, the vector is an expression vector. In another embodiment, the expression vector includes a promoter operably linked to one or more of the polynucleotides of the invention. In another embodiment, a cell comprises a vector that includes a polynucleotide of the invention.

[0055] In another aspect, the invention provides compositions of compounds and methods of producing such compounds. For example, compounds include, e.g., an unnatural amino acid (such as *p*-(propargyloxy)-phenyalanine (e.g., 1 in **Figure 11**), azido dyes (such as shown in chemical structure 4 and chemical structure 6), an alkynyl polyethylene glycol (e.g., as shown in chemical structure 7), where n is an integer between, e.g., 50 and 10,000, 75 and 5,000, 100 and 2,000, 100 and 1,000, etc., and the like. In embodiment of the invention, the alkynyl polyethylene glycol has a molecular weight of, e.g., about 5,000 to about 100,000 Da, about 20,000 to about 50,000 Da, about 20,000 to about 10,000 Da (e.g., 20,000 Da).

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Various compositions comprising these compounds, e.g., with proteins and cells, are also provided. In one aspect, the composition that includes the *p*-(propargyloxy)-phenyalanine unnatural amino acid, further includes an orthogonal tRNA. The unnatural amino acid can be bonded (e.g., covalently) to the orthogonal tRNA, e.g., covalently bonded to the orthogonal tRNA though an amino-acyl bond, covalently bonded to a 3'OH or a 2'OH of a terminal ribose sugar of the orthogonal tRNA, etc.

[0057] In one aspect of the invention, a protein comprising an azido dye (e.g., of chemical structure 4 or chemical structure 6), further includes at least one unnatural amino acid (e.g., an alkynyl amino acid), where the azido dye is attached to the unnatural amino acid through a [3+2] cycloaddition.

In one embodiment, a protein comprises the alkynyl polyethylene glycol of chemical structure 7. In another embodiment, the composition further includes at least one unnatural amino acid (e.g., an azido amino acid), wherein the alkynyl polyethylene glycol is attached to an unnatural amino acid through a [3+2] cycloaddition.

[0059] Methods for synthesizing various compounds are included in the invention.

For example, a method for synthesizing a p-(propargyloxy)phenyalanine compound is provided. For example, the method comprises (a) suspending N-tert-butoxycarbonyl-tyrosine and K<sub>2</sub>CO<sub>3</sub> in anhydrous DMF; (b) adding propargyl bromide to the reaction mixture of (a) and alkylating the hydroxyl and the carboxyl group, resulting in an protected intermediate compound having the structure:

and (c) mixing the protected intermediate compound with anhydrous HCl in MeOH and deprotecting the amine moiety, thereby synthesizing the *p*-(propargyloxy)phenyalanine compound. In one embodiment, the method further comprises (d) dissolving the p-(propargyloxy)phenylalanine HCl in aqueous NaOH and MeOH and stirring it at room temperature; (e) adjusting the pH of to pH 7; and (f) precipitating the p-(propargyloxy)phenylalanine compound.

[0060] Methods for synthesizing azido dyes are also provided. For example, a method comprises: (a) providing a dye compound comprising a sulfonyl halide moiety; (b) warming the dye compound to room temperature in the presence of 3-azidopropylamine and triethylamine and coupling an amine moiety of the 3-azidopropylamine to the halide position of the dye compound, thereby synthesizing the azido dye. In one embodiment, the dye compound comprises dansyl chloride, and the azido dye comprises the composition of chemical structure 4. In one aspect, the method further comprises purifying the azido dye from the reaction mixture.

providing an amine-containing dye compound; (b) combining the amine-containing dye compound with a carbodiimide and 4-(3-azidopropylcarbamoyl)-butyric acid in a suitable solvent, and coupling a carbonyl group of the acid to the amine moiety of the dye compound, thereby synthesizing the azido dye. In one embodiment, the carbodiimine comprises 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochlŏride (EDCI). In one aspect, the amine-containing dye comprises fluoresceinamine, and the suitable solvent comprises pyridine. For example, the amine-containing dye comprises fluoresceinamine and the azido dye comprises the composition of chemical structure 6. In one embodiment, the method further comprises (c) precipitating the azido dye; (d) washing the precipitate with HCl; (e) dissolving the washed precipitate in EtOAc; and (f) precipitating the azido dye in hexanes.

[0062] Methods for synthesizing a propargyl amide polyethylene glycol are also provided. For example, the method comprises reacting propargylamine with polyethylene glycol (PEG)-hydroxysuccinimide ester in an organic solvent (e.g., CH<sub>2</sub>Cl<sub>2</sub>) at room temperature, resulting in the propargyl amide polyethylene glycol of chemical structure 7.

In one embodiment, the method further comprises precipitating the propargylamide polyethylene glycol using ethyl acetate. In one aspect, the method further includes recrystallizing the propargylamide polyethylene glycol in methanol; and drying the product under a vacuum.

[0063] Kits are also a feature of the invention. For example, a kit for producing a protein that comprises at least one unnatural amino acid in a cell is provided, where the kit includes a container containing a polynucleotide sequence encoding an O-tRNA or an O-tRNA, and a polynucleotide sequence encoding an O-RS or an O-RS. In one embodiment, the kit further includes at least one unnatural amino acid. In another embodiment, the kit further comprises instructional materials for producing the protein.

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# **DEFINITIONS**

[0064] Before describing the present invention in detail, it is to be understood that this invention is not limited to particular devices or biological systems, which can, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to be limiting. As used in this specification and the appended claims, the singular forms "a", "an" and "the" include plural referents unless the content clearly dictates otherwise. Thus, for example, reference to "a cell" includes a combination of two or more cells; reference to "bacteria" includes mixtures of bacteria, and the like.

[0065] Unless otherwise defined herein or below in the remainder of the specification, all technical and scientific terms used herein have the same meaning as commonly understood by those of ordinary skill in the art to which the invention belongs.

Homologous: Proteins and/or protein sequences are "homologous" when they are derived, naturally or artificially, from a common ancestral protein or protein sequence. Similarly, nucleic acids and/or nucleic acid sequences are homologous when they are derived, naturally or artificially, from a common ancestral nucleic acid or nucleic acid sequence. For example, any naturally occurring nucleic acid can be modified by any available mutagenesis method to include one or more selector codon. When expressed, this mutagenized nucleic acid encodes a polypeptide comprising one or more unnatural amino acid. The mutation process can, of course, additionally alter one or more standard codon, thereby changing one or more standard amino acid in the resulting mutant protein, as well.

Homology is generally inferred from sequence similarity between two or more nucleic acids or proteins (or sequences thereof). The precise percentage of similarity between sequences that is useful in establishing homology varies with the nucleic acid and protein at issue, but as little as 25% sequence similarity is routinely used to establish homology. Higher levels of sequence similarity, e.g., 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% or more, can also be used to establish homology. Methods for determining sequence similarity percentages (e.g., BLASTP and BLASTN using default parameters) are described herein and are generally available.

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Orthogonal: As used herein, the term "orthogonal" refers to a molecule (e.g., [0067] an orthogonal tRNA (O-tRNA) and/or an orthogonal aminoacyl tRNA synthetase (O-RS)) that functions with endogenous components of a cell with reduced efficiency as compared to a corresponding molecule that is endogenous to the cell or translation system, or that fails to function with endogenous components of the cell. In the context of tRNAs and aminoacyl-tRNA synthetases, orthogonal refers to an inability or reduced efficiency, e.g., less than 20 % efficient, less than 10 % efficient, less than 5 % efficient, or less than 1% efficient, of an orthogonal tRNA to function with an endogenous tRNA synthetase compared to an endogenous tRNA to function with the endogenous tRNA synthetase, or of an orthogonal aminoacyl-tRNA synthetase to function with an endogenous tRNA compared to an endogenous tRNA synthetase to function with the endogenous tRNA. The orthogonal molecule lacks a functional endogenous complementary molecule in the cell. For example, an orthogonal tRNA in a cell is aminoacylated by any endogenous RS of the cell with reduced or even zero efficiency, when compared to aminoacylation of an endogenous tRNA by the endogenous RS. In another example, an orthogonal RS aminoacylates any endogenous tRNA in a cell of interest with reduced or even zero efficiency, as compared to aminoacylation of the endogenous tRNA by an endogenous RS. A second orthogonal molecule can be introduced into the cell that functions with the first orthogonal molecule. For example, an orthogonal tRNA/RS pair includes introduced complementary components that function together in the cell with an efficiency (e.g., 50% efficiency, 60% efficiency, 70% efficiency, 75% efficiency, 80% efficiency, 90% efficiency, 95% efficiency, or 99% or more efficiency) to that of a corresponding tRNA/RS endogenous pair.

[0068] Complementary: The term "complementary" refers to components of an orthogonal pair, O-tRNA and O-RS that can function together, e.g., where the O-RS aminoacylates the O-tRNA.

[0069] Preferentially aminoacylates: The term "preferentially aminoacylates" refers to an efficiency, e.g., 70 % efficient, 75 % efficient, 85% efficient, 90% efficient, 95 % efficient, or 99% or more efficient, at which an O-RS aminoacylates an O-tRNA with an unnatural amino acid as compared to the O-RS aminoacylating a naturally occurring tRNA or a starting material used to generate the O-tRNA. The unnatural amino acid is incorporated into a growing polypeptide chain with high fidelity, e.g., at greater than 75% efficiency for a given selector codon, at greater than about 80% efficiency for a given selector codon, at greater than about 95% efficiency for a given selector codon, or at greater than about 99% or more efficiency for a given selector codon.

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[0070] Selector codon: The term "selector codon" refers to codons recognized by the O-tRNA in the translation process and not recognized by an endogenous tRNA. The O-tRNA anticodon loop recognizes the selector codon on the mRNA and incorporates its amino acid, e.g., an unnatural amino acid, at this site in the polypeptide. Selector codons can include, e.g., nonsense codons, such as, stop codons, e.g., amber, ochre, and opal codons; four or more base codons; rare codons; codons derived from natural or unnatural base pairs and/or the like.

[0071] Suppressor tRNA: A suppressor tRNA is a tRNA that alters the reading of a messenger RNA (mRNA) in a given translation system, e.g., by providing a mechanism for incorporating an amino acid into a polypeptide chain in response to a selector codon. For example, a suppressor tRNA can read through, e.g., a stop codon, a four base codon, a rare codon, and/or the like.

[0072] Recyclable tRNA: The term "recyclable tRNA" refers to a tRNA that is aminoacylated and can be repeatedly reaminoacylated with an amino acid (e.g., an unnatural amino acid) for the incorporation of the amino acid (e.g., the unnatural amino acid) into one or more polypeptide chains during translation.

Translation system: The term "translation system" refers to the collective set of components that incorporate a naturally occurring amino acid into a growing polypeptide chain (protein). Components of a translation system can include, e.g., ribosomes, tRNAs, synthetases, mRNA, amino acids, and the like. The components of the invention (e.g., ORS, OtRNAs, unnatural amino acids, etc.) can be added to an in vitro or in vivo translation

system, e.g., a eukaryotic cell, e.g., a yeast cell, a mammalian cell, a plant cell, an algae cell, a fungus cell, an insect cell, and/or the like.

[0074] <u>Unnatural amino acid</u>: As used herein, the term "unnatural amino acid" refers to any amino acid, modified amino acid, and/or amino acid analogue that is not one of the 20 common naturally occurring amino acids, seleno cysteine or pyrrolysine.

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[0075] <u>Derived from</u>: As used herein, the term "derived from" refers to a component that is isolated from or made using information from a specified molecule or organism.

[0076] <u>Inactive RS</u>: As used herein, the term "inactive RS" refers to a synthetase that has been mutated so that it no longer can aminoacylate its natural cognate tRNA with an amino acid.

[0077] <u>Positive selection or screening marker</u>: As used herein, the term "positive selection or screening marker" refers to a marker that when present, e.g., expressed, activated or the like, results in identification of a cell with the positive selection marker from those without the positive selection marker.

15 [0078] Negative selection or screening marker: As used herein, the term "negative selection or screening marker" refers to a marker that when present, e.g., expressed, activated or the like, allows identification of a cell that does not possess the desired property (e.g., as compared to a cell that does possess the desired property).

[0079] Reporter: As used herein, the term "reporter" refers to a component that can be used to select target components of a system of interest. For example, a reporter can include a fluorescent screening marker (e.g., green fluorescent protein), a luminescent marker (e.g., a firefly luciferase protein), an affinity based screening marker, or selectable marker genes such as his3, ura3, leu2, lys2, lacZ, β-gal/lacZ (β-galactosidase), Adh (alcohol dehydrogenase), or the like.

25 [0080] <u>Eukaryote</u>: As used herein, the term "eukaryote" refers to organisms belonging to the phylogenetic domain Eucarya such as animals (e.g., mammals, insects, reptiles, birds, etc.), ciliates, plants (e.g., monocots, dicots, algae, etc.), fungi, yeasts, flagellates, microsporidia, protists, etc.

[0081] Non-eukaryote: As used herein, the term "non-eukaryote" refers to non-eukaryotic organisms. For example, a non-eukaryotic organism can belong to the Eubacteria (e.g., Escherichia coli, Thermus thermophilus, Bacillus stearothermophilus, etc.)

phylogenetic domain, or the Archaea (e.g., Methanococcus jannaschii, Methanobacterium thermoautotrophicum, Halobacterium such as Haloferax volcanii and Halobacterium species NRC-1, Archaeoglobus fulgidus, Pyrococcus furiosus, Pyrococcus horikoshii, Aeuropyrum pernix, etc.) phylogenetic domain.

Antibody: The term "antibody," as used herein, includes, but is not limited to a polypeptide substantially encoded by an immunoglobulin gene or immunoglobulin genes, or fragments thereof, which specifically bind and recognize an analyte (antigen). Examples include polyclonal, monoclonal, chimeric, and single chain antibodies, and the like. Fragments of immunoglobulins, including Fab fragments and fragments produced by an expression library, including phage display, are also included in the term "antibody" as used herein. See, e.g., Paul, Fundamental Immunology, 4th Ed., 1999, Raven Press, New York, for antibody structure and terminology.

Conservative variant: The term "conservative variant" refers to a translation component, e.g., a conservative variant O-tRNA or a conservative variant O-RS, that functionally performs like the component from which the conservative variant is based, e.g., an O-tRNA or O-RS, but has variations in the sequence. For example, an O-RS will aminoacylate a complementary O-tRNA or a conservative variant O-tRNA with an unnatural amino acid, although the O-tRNA and the conservative variant O-tRNA do not have the same sequence. The conservative variant can have, e.g., one variation, two variations, three variations, four variations, or five or more variations in sequence, as long as the conservative variant is complementary to the corresponding O-tRNA or O-RS.

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Selection or screening agent: As used herein, the term "selection or screening agent" refers to an agent that, when present, allows for a selection/screening of certain components from a population. For example, a selection or screening agent includes, but is not limited to, e.g., a nutrient, an antibiotic, a wavelength of light, an antibody, an expressed polynucleotide (e.g., a transcriptional modulator protein), or the like. The selection agent can be varied, e.g., by concentration, intensity, etc.

[0085] <u>Detectable substance</u>: The term "detectable substance," as used herein, refers to an agent that, when activated, altered, expressed or the like, allows for the selection/screening of certain components from a population. For example, the detectable substance can be a chemical agent, e.g., 5-fluroorotic acid (5-FOA), which under certain

conditions, e.g., expression of a URA3 reporter, becomes detectable, e.g., a toxic product that kills cells that express the URA3 reporter.

# BRIEF DESCRIPTION OF THE DRAWINGS

- [0086] Figure 1, Panels A, B and C schematically illustrates a general positive and negative selection scheme for expanding the genetic code of a eukaryotic cell, e.g., S. cerevisiae. Panel A schematically illustrates the activated transcription of reporter genes, which are is driven by amber suppression of TAG codons in GALA. The DNA binding domain is indicated by the striped box and the major and cryptic activation domains are indicated in the hatched box. Panel B illustrates examples of reporter genes, e.g., HIS3, LacZ, URA3 in MaV203. Panel C schematically illustrates plasmids that can be used in the selection scheme, e.g., pEcTyrRS/tRNA<sub>CUA</sub> and pGADGAL4xxTAG.
  - [0087] Figure 2 illustrates the EcTyrRS and tRNA<sub>CUA</sub> dependent phenotypes of 1<sup>st</sup> generation GAL4 reporters on selective media. DB-AD is a fusion between the GAL4 DNA binding domain and activation domain. DB-TAG-AD has a TAG codon replacing a tyrosine codon in the synthetic linker between DB and AD. A5 is an inactive version of EcTyrRS in which 5 residues in the active site have been mutated to alanine.

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- [0088] Figure 3, Panel A and B illustrate EcTyrRS and tRNA<sub>CUA</sub> dependent phenotypes of 2nd generation GAL4 reporters on selective media. The DNA binding domain is indicated by the striped box and the major and cryptic activation domains are indicated in the hatched box. Panel A illustrates constructs each with single amino acid mutation in GAL4. Panel B illustrates constructs each with two amino acid mutations in GAL4.
- [0089] Figure 4 Panels A, B and C illustrate pGADGAL4 (T44TAG, R110TAG) with and without EcTyrRS and various reporters in MaV203. Panel A shows the results in the presence of X-gal, -Ura, or -Leu, -Trp. Panel B shows the results in the presence of varying concentrations of 3-AT. Panel C shows the results in the present of varying percentages of 5-FOA.
- [0090] Figure 5 Panels A and B illustrate ONPG hydrolysis with various GAL4 mutants, e.g., where residues T44 (A) and R110 (B) are permissive sites. Panel A

  30 illustrates the ONPG hydrolysis measurement with various types of mutations at the T44 site. Panel B illustrates the ONPG hydrolysis measurement with various types of mutations at the R110 site. 'GAL4' is MaV203 transformed with pCL1 and was offscale ~ 600 ONPG

hydrolysis units. 'None' is MaV203 transformed with plasmids encoding the GAL4 DB and GAL4 AD separately.

[0091] Figure 6 shows the selection of active EcTyrRS clones. MaV203 containing a 1:10 mixture of pEcTyrRS-tRNA<sub>CUA</sub>: pA5-tRNA<sub>CUA</sub> were plated at a 10<sup>3</sup> dilution on (-Leu, -Trp) plates (left) or (-Leu, -Trp,-His + 50 mM 3-AT) plates (right) and processed using XGAL overlay.

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[0092] Figure 7, Panels A and B. Panel A illustrates a stereoview of the active site of *B. Stearothermophilus* tyrosyl-tRNA synthetase with bound tyrosine. The mutated residues are shown and correspond to residues from *E. coli* tyrosyl-tRNA synthetase Tyr<sup>37</sup> (*B. Stearothermophilus* TyrRS residue Tyr<sup>34</sup>), Asn<sup>126</sup> (Asn<sup>123</sup>), Asp<sup>182</sup> (Asp<sup>176</sup>), Phe<sup>183</sup> (Phe<sup>177</sup>), and Leu<sup>186</sup> (Leu<sup>180</sup>). Panel B illustrates the structural formulae of examples of unnatural amino acids (from left to right) *p*-acetyl-L-phenylalanine (1), *p*-benzoyl-L-phenylalanine (2), *p*-azido-L-phenylalanine (3), O-methyl-L-tyrosine (4), and *p*-iodo-L-tyrosine (5).

15 [0093] Figure 8, Panels A, B, C and D. Panel A illustrates vectors and reporter constructs that can be used in selection/screening for orthogonal tRNAs, orthogonal aminoacyl synthetases or pairs of orthogonal tRNA/RS in eukaryotic cells. Panel B illustrates phenotypes of yeast harboring GAL4 responsive HIS3, URA3 and lacZ responsive reporters in response to active (TyrRS) or inactive (A5RS) aminoacyl-tRNA synthetases on selective media. Panel C illustrates an example of a selection scheme used to select mutant synthetases that encode additional amino acids in a eukaryotic cell, e.g., S. cerevisiae, where UAA is an unnatural amino acid. Panel D illustrates phenotypes of yeast isolated from a selection with p-acetyl-L-phenylalanine.

[0094] Figure 9 illustrates protein expression of human superoxide dismutase (hSOD) (33TAG)HIS in *S. cerevisiae* genetically encoding unnatural amino acids as indicated in Figure 7, Panel B.

[0095] Figure 10, Panels A-H illustrate tandem mass spectrum of the tryptic peptide VY\*GSIK (SEQ ID NO:87) containing the unnatural amino acids (denoted Y\*) as indicated in Figure 7, Panel B. Panel A illustrates tandem mass spectrum of tryptic peptide with unnatural amino acid p-acetyl-L-phenylalanine (1). Panel B illustrates tandem mass spectrum of tryptic peptide with unnatural amino acid p-benzoyl-L-phenylalanine (2). Panel C illustrates tandem mass spectrum of tryptic peptide with unnatural amino acid p-

- azido-L-phenylalanine (3). Panel D illustrates tandem mass spectrum of tryptic peptide with unnatural amino acid O-methyl-L-tyrosine (4). Panel E illustrates tandem mass spectrum of tryptic peptide with unnatural amino acid p-iodo-L-tyrosine (5). Panel F illustrates tandem mass spectrum of tryptic peptide with amino acid trytophan (W) at position Y\*. Panel G illustrates tandem mass spectrum of tryptic peptide with amino acid tyrosine (Y) at position Y\*. Panel H illustrates tandem mass spectrum of tryptic peptide with amino acid leucine (L) at position Y\*.
- [0096] Figure 11 illustrates examples of two unnatural amino acids (1) para-propargyloxyophenylalanine and (2) para-azidophenylalanine.

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- 10 [0097] Figure 12, Panels A, B and C illustrate SOD expression in the presence or absence of unnatural amino acids 1 and 2 indicated in Figure 11. Panel A illustrates Gelcode Blue stain experiment. Panel B illustrates a western blot with an anti-SOD antibody. Panel C illustrates a western blot with anti-6xHis antibody.
- [0098] Figure 13, Panels A, B and C illustrate protein labeling by [3+2] cycyloaddition. Panel A illustrates synthesized dye labels 3-6. Panel B illustrates the reaction between the SOD and the dye. Panel C illustrates in-gel fluorescence scanning and Gelcode Blue staining.
  - [0099] Figure 14 illustrates growth of eukaryotic cells, e.g., S. cerevisiae cells, transformed with synthetase mutants in the absence or presence of 1 or 2 as indicated in Figure 11 on SD media lacking uracil.
  - [0100] Figure 15, Panels A and B, illustrate tandem mass spectrum of the tryptic peptide VY\*GSIK (SEQ ID NO:87) containing the azide (Az) (Panel A) or alkyne (Al) (Panel B) unnatural amino acids in position Y\* are shown with their expected fragment ion masses. Arrow indicates observed b (blue) and y (red) ions series for each peptide.
- 25 **[0101]** Figure 16 schematically illustrates the in vivo incorporation of an unnatural amino acid, e.g., *para*-propargyloxyophenylalanine, into a growing polypeptide chain and the bioconjugation with small organic molecules by a [3+2]-cycloaddition reaction through this unnatural amino acid.
- [0102] Figure 17, Panels A, B and C illustrate PEGylation of a protein comprising an unnatural amino acid using a [3+2] cycloaddition. Panel A illustrates the reaction of a propargyl amide PEG with a protein comprising an azido amino acid (e.g., N<sub>3</sub>-SOD) in the

presence of Cu(I) and phosphate buffer (PB). Panel B illustrates the PEGylation of the protein by gel analysis. Panel C illustrates the synthesis of the propargyl amide PEG.

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#### **DETAILED DESCRIPTION**

[0103] The ability to genetically modify the structures of proteins directly in eukaryotic cells, beyond the chemical constraints imposed by the genetic code, would provides a powerful molecular tool to both probe and manipulate cellular processes. The invention provides translational components that expand the number of genetically encoded amino acids in eukaryotic cells. These include tRNAs (e.g., orthogonal tRNAs (O-tRNAs)), aminoacyl-tRNA synthetases (e.g., orthogonal synthetase (O-RS)), pairs of O-tRNA/O-RSs, and unnatural amino acids.

[0104] Typically, O-tRNAs of the invention are expressed and processed efficiently, and function in translation in a eukaryotic cell, but are not significantly aminoacylated by the host's aminoacyl-tRNA synthetases. In response to a selector codon, an O-tRNA of the invention delivers an unnatural amino acid, which does not encode any of the common twenty amino acids, to a growing polypeptide chain during mRNA translation.

[0105] An O-RS of the invention preferentially aminoacylates an O-tRNA of the invention with an unnatural amino acid in a eukaryotic cell, but does not aminoacylate any of the cytoplasmic host's tRNAs. Moreover, the specificity of an aminoacyl-tRNA synthetase of the invention provides acceptance of an unnatural amino acid while excluding any endogenous amino acids. Polypeptides that include amino acid sequences of example O-RSs, or portions thereof, are also a feature of the invention. In addition, polynucleotides that encode translational components, O-tRNAs, O-RSs and portions thereof, are features of the invention.

[0106] The invention also provides methods of producing the desired translational components, e.g., O-RS, and or an orthogonal pair (orthogonal tRNA and orthogonal aminoacyl-tRNA synthetase), that utilizes an unnatural amino acid for use in a eukaryotic cell (and translational components produced by such methods). For example, a tyrosyl-tRNA synthetase/tRNA<sub>CUA</sub> pair from *E. coli* is an O-tRNA/O-RS pair of the invention. In addition, the invention also features methods of selecting/screening translational components in one eukaryotic cell, and once selected/screened, using those components in a different eukaryotic cell (a eukaryotic cell that was not used for selection/screening). For example, the selection/screening methods to produce the translation components for

eukaryotic cells can be done in yeast, e.g., *Saccharomyces cerevisiae*, and then those selected components can be used in another eukaryotic cell, e.g., another yeast cell, a mammalian cell, an insect cell, a plant cell, a fungus cell, etc.

[0107] The invention further provides methods for producing a protein in a eukaryotic cell, where the protein comprises an unnatural amino acid. The protein is produced using the translation components of the invention. The invention also provides proteins (and proteins produced by the methods of the invention), which include unnatural amino acids. The protein or polypeptide of interest can also include a post-translational modification, e.g., that is added through a [3+2] cycloaddition, or a nucleophilic-electrophilic reaction, that is not made by a prokaryotic cell, etc. In certain embodiments, methods of producing a transcriptional modulator protein with an unnatural amino acid (and proteins produced by such methods) are also included in the invention. Compositions, which include proteins that include an unnatural amino acid is also a feature of the invention.

15 [0108] Kits for producing a protein or polypeptide with an unnatural amino acid are also a feature of the invention.

# ORTHOGONAL AMINOACYL-tRNA SYNTHETASES (O-RS)

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[0109] In order to specifically incorporate an unnatural amino acid in to a protein or polypeptide of interest, in a eukaryotic cell, the substrate specificity of the synthetase is altered so that only the desired unnatural amino acid, but not any of the common 20 amino acids are charged to the tRNA. If the orthogonal synthetase is promiscuous, it will result in mutant proteins with a mixture of natural and unnatural amino acids at the target position. The invention provides compositions of, and methods of, producing orthogonal aminoacyltRNA synthetases that have modified substrate specificity for a specific unnatural amino acid.

[0110] A eukaryotic cell that includes an orthogonal aminoacyl-tRNA synthetase (O-RS) is a feature of the invention. The O-RS preferentially aminoacylates an orthogonal tRNA (O-tRNA) with an unnatural amino acid in the eukaryotic cell. In certain embodiments, the O-RS utilizes more than one unnatural amino acid, e.g., two or more, three or more, etc. Thus, an O-RS of the invention can have the capability to preferentially aminoacylate an O-tRNA with different unnatural amino acids. This allows an additional level of control by selecting which unnatural amino acid or combination of unnatural amino

acids are put with the cell and/or by selecting the different amounts of unnatural amino acids that are put with the cell for their incorporation.

[0111] An O-RS of the invention optionally has one or more improved or enhanced enzymatic properties for the unnatural amino acid as compared to a natural amino acid. These properties include, e.g., higher Km, lower Km, higher kcat, lower kcat, lower kcat/km, higher kcat/km, etc., for the unnatural amino acid, as compared to a naturally occurring amino acid, e.g., one of the 20 known common amino acids.

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[0112] Optionally, the O-RS can be provided to the eukaryotic cell by a polypeptide that includes an O-RS and/or by a polynucleotide that encodes an O-RS or a portion thereof. For example, an O-RS, or a portion thereof, is encoded by a polynucleotide sequence as set forth in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), or a complementary polynucleotide sequence thereof. In another example, an O-RS comprises an amino acid sequence as set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, or a conservative variation thereof. See, e.g., Tables 5, 6 and 8, and Example 6 herein for sequences of exemplary O-RS molecules.

An O-RS can also comprise an amino acid sequence that is, e.g., at least [0113] 90%, at least 95%, at least 98%, at least 99%, or even at least 99.5 % identical to that of a naturally occurring tyrosyl aminoacyl-tRNA synthetase (TyrRS) (e.g., as set forth in SEQ ID NO.:2) and comprises two or more amino acids of group A-E. Group A includes valine, isoleucine, leucine, glycine, serine, alanine, or threonine at a position corresponding to Tyr37 of E. coli TyrRS; group B includes aspartate at a position corresponding to Asn126 of E. coli TyrRS; group C includes threonine, serine, arginine, asparagine or glycine at a position corresponding to Asp182 of E. coli TyrRS; group D includes methionine, alanine, valine, or tyrosine at a position corresponding to Phe183 of E. coli TyrRS; and, group E includes serine, methionine, valine, cysteine, threonine, or alanine at a position corresponding to Leu186 of E. coli TyrRS. Any subset of combinations of these groups are a feature of the invention. For example, in one embodiment, the O-RS has two or more amino acids selected from valine, isoleucine, leucine, or threonine occurs at a position corresponding to Tyr37 of E. coli TyrRS; threonine, serine, arginine, or glycine at a position corresponding to Asp182 of E. coli TyrRS; methionine, or tyrosine at a position corresponding to Phe183 of E. coli TyrRS; and, serine, or alanine at a position corresponding to Leu186 of E. coli TyrRS. In another embodiment, the O-RS includes two

more more amino acids selected from glycine, serine, or alanine at a position corresponding to Tyr37 of *E. coli* TyrRS, aspartate at a position corresponding to Asn126 of E. coli TyrRS, asparagine at a position corresponding to Asp182 of *E. coli* TyrRS, alanine, or valine, at a position corresponding to Phe183 of *E. coli* TyrRS, and/or methionine, valine, cysteine, or threonine, at a position corresponding to Leu186 of *E. coli* TyrRS. *See also, e.g.*, Table 4, Table 6 and Table 8, herein.

[0114] Besides the O-RS, a eukaryotic cell of the invention can include additional components, e.g., an unnatural amino acid(s). The eukaryotic cell also includes an orthogonal tRNA (O-tRNA) (e.g., derived from a non-eukaryotic organism, such as *Escherichia coli*, *Bacillus stearothermophilus*, and/or the like), where the O-tRNA recognizes a selector codon and is preferentially aminoacylated with the unnatural amino acid by the O-RS. A nucleic acid that comprises a polynucleotide that encodes a polypeptide of interest, wherein the polynucleotide comprises a selector codon that is recognized by the O-tRNA, or a combination of one or more of these, can also be present in the cell.

In one aspect, the O-tRNA mediates the incorporation of the unnatural amino acid into a protein with, e.g., at least 45%, at least 50%, at least 60%, at least 75%, at least 80%, at least 90%, at least 95%, or 99% or the efficiency of as a tRNA that comprises or is processed from a polynucleotide sequence as set forth in SEQ ID NO.: 65. In another aspect, the O-tRNA comprises SEQ ID NO.:65, and the O-RS comprises a polypeptide sequence set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, and/or a conservative variation thereof. *See also*, e.g., Table 5 and Example 6, herein, for sequences of exemplary O-RS and O-tRNA molecules.

[0116] In one example, a eukaryotic cell comprises an orthogonal aminoacyl-tRNA synthetase (O-RS), an orthogonal tRNA (O-tRNA), an unnatural amino acid, and a nucleic acid that comprises a polynucleotide that encodes a polypeptide of interest, which polynucleotide comprises a selector codon that is recognized by the O-tRNA. The O-RS preferentially aminoacylates the orthogonal tRNA (O-tRNA) with the unnatural amino acid in the eukaryotic cell, and the cell produces the polypeptide of interest in the absence of the unnatural amino acid with a yield that is, e.g., less than 30%, less than 20%, less than 15%, less than 5%, less than 2.5%, etc., of the yield of the polypeptide in the presence of the unnatural amino acid.

[0117] Methods for producing an O-RS, which are a feature of the invention, optionally include generating a pool of mutant synthetases from the framework of a wild-type synthetase, and then selecting for mutated RSs based on their specificity for an unnatural amino acid relative to the common twenty amino acids. To isolate such a synthetase, the selection methods of the are: (i) sensitive, as the activity of desired synthetases from the initial rounds can be low and the population small; (ii) "tunable", since it is desirable to vary the selection stringency at different selection rounds; and, (iii) general, so that the methods can be used for different unnatural amino acids.

that preferentially aminoacylates an orthogonal tRNA with an unnatural amino acid in a eukaryotic cell typically include applying a combination of a positive selection followed by a negative selection. In the positive selection, suppression of the selector codon introduced at nonessential position(s) of a positive marker allows the eukaryotic cells to survive under positive selection pressure. In the presence of unnatural amino acids, survivors thus encode active synthetases charging the orthogonal suppressor tRNA with an unnatural amino acid. In the negative selection, suppression of a selector codon introduced at nonessential position(s) of a negative marker removes synthetases with natural amino acid specificities. Survivors of the negative and positive selection encode synthetases that aminoacylate (charge) the orthogonal suppressor tRNA with unnatural amino acids only (or at least preferentially).

[0119] For example, the method includes: (a) subjecting to positive selection, in the presence of an unnatural amino acid, a population of eukaryotic cells of a first species, where the eukaryotic cells each comprise: i) a member of a library of aminoacyl-tRNA synthetases (RSs), ii) an orthogonal tRNA (O-tRNA), iii) a polynucleotide that encodes a positive selection marker, and iv) a polynucleotide that encodes a negative selection marker; wherein cells that survive the positive selection comprise an active RS that aminoacylates the orthogonal tRNA (O-tRNA) in the presence of an unnatural amino acid; and, (b) subjecting the cells that survive the positive selection to negative selection in the absence of the unnatural amino acid to eliminate active RSs that aminoacylate the O-tRNA with a natural amino acid, thereby providing the O-RS that preferentially aminoacylates the O-tRNA with the unnatural amino acid.

[0120]The positive selection marker can be any of a variety of molecules. In one embodiment, the positive selection marker is a product that provides a nutritional supplement for growth and the selection is performed on a medium that lacks the nutritional supplement. Examples of polynucleotides that encode positive selection markers include, 5 but are not limited to, e.g., a reporter gene based on complementing the amino acid auxotrophy of a cell, a his3 gene (e.g., where the his3 gene encodes an imidazole glycerol phosphate dehydratase, detected by providing 3-aminotriazole (3-AT)), ura3 gene, leu2 gene, lys2 gene, lacZ gene, adh gene, etc. See, e.g., G.M. Kishore, & D.M. Shah, (1988), Amino acid biosynthesis inhibitors as herbicides, Annual Review of Biochemistry 57:627-10 663. In one embodiment, lacZ production is detected by ortho-nitrophenyl-β-Dgalactopyranoside (ONPG) hydrolysis. See, e.g., I.G. Serebriiskii, & E.A. Golemis, (2000), Uses of lacZ to study gene function: evaluation of beta-galactosidase assays employed in the yeast two-hybrid system, Analytical Biochemistry 285:1-15. Additional positive selection markers include, e.g., luciferase, green fluorescent protein (GFP), YFP, EGFP, 15 RFP, the product of an antibiotic resistant gene (e.g., chloramphenicol acetyltransferase (CAT)), a transcriptional modulator protein (e.g., GAL4), etc. Optionally, a polynucleotide that encodes a positive selection marker comprises a selector codon.

linked to a response element. An additional polynucleotide that encodes a transcriptional modulator protein that modulates transcription from the response element, and comprises at least one selector codon, can also be present. The incorporation of the unnatural amino acid into the transcriptional modulator protein by the O-tRNA aminoacylated with the unnatural amino acid results in transcription of the polynucleotide (e.g., reporter gene) encoding the positive selection marker. For example, see **Figure 1A**. Optionally, the selector codon is located in or substantially near a portion of the polynucleotide that encodes a DNA binding domain of the transcriptional modulator protein.

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[0122] A polynucleotide that encodes the negative selection marker can also be operably linked to a response element from which transcription is mediated by the transcriptional modulator protein. See, e.g., A.J. DeMaggio, et al., (2000), The yeast splithybrid system, Method Enzymol. 328:128-137; H.M. Shih, et al., (1996), A positive genetic selection for disrupting protein-protein interactions: identification of CREB mutations that prevent association with the coactivator CBP, Proc. Natl. Acad. Sci. U. S. A. 93:13896-13901; M. Vidal, et al., (1996), Genetic characterization of a mammalian protein-protein

interaction domain by using a yeast reverse two-hybrid system. [comment], Proc. Natl.

Acad. Sci. U. S. A. 93:10321-10326; and, M. Vidal, et al., (1996), Reverse two-hybrid and one-hybrid systems to detect dissociation of protein-protein and DNA-protein interactions. [comment], Proc. Natl. Acad. Sci. U. S. A. 93:10315-10320. The incorporation of a natural amino acid into the transcriptional modulator protein by the O-tRNA aminoacylated with a natural amino acid results in transcription of the negative selection marker. Optionally, the negative selection marker comprises a selector codon. In one embodiment, the positive selection marker and/or negative selection marker of the invention can comprise at least two selector codons, which each or both can comprise at least two different selector codons or at least two of the same selector codons.

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- [0123] The transcriptional modulator protein is a molecule that binds (directly or indirectly) to a nucleic acid sequence (e.g., a response element) and modulates transcription of a sequence that is operably linked to the response element. A transcriptional modulator protein can be a transcriptional activator protein (e.g., GAL4, nuclear hormone receptors, AP1, CREB, LEF/tcf family members, SMADs, VP16, SP1, etc.), a transcriptional repressor protein (e.g., nuclear hormone receptors, Groucho/tle family, Engrailed family, etc), or a protein that can have both activities depending on the environment (e.g., LEF/tcf, homobox proteins, etc.). A response element is typically a nucleic acid sequence that is recognized by the transcriptional modulator protein or an additional agent that acts in concert with the transcriptional modulator protein.
- [0124] Another example of a transcriptional modulator protein is the transcriptional activator protein, GAL4 (see e.g., Figure 1A). See, e.g., A. Laughon, et al., (1984), Identification of two proteins encoded by the Saccharomyces cerevisiae GALA gene, Molecular & Cellular Biology 4:268-275; A. Laughon, & R.F. Gesteland, (1984), Primary structure of the Saccharomyces cerevisiae GALA gene, Molecular & Cellular Biology 25 4:260-267; L. Keegan, et al., (1986), Separation of DNA binding from the transcriptionactivating function of a eukaryotic regulatory protein, Science 231:699-704; and, M. Ptashne, (1988), How eukaryotic transcriptional activators work, Nature 335:683-689. The N-terminal 147 amino acids of this 881 amino acid protein form a DNA binding domain (DBD) that binds DNA sequence specifically. See, e.g., M. Carey, et al., (1989), An amino-30 terminal fragment of GAL4 binds DNA as a dimer, J. Mol. Biol. 209:423-432; and, E. Giniger, et al., (1985), Specific DNA binding of GALA, a positive regulatory protein of yeast, Cell 40:767-774. The DBD is linked, by an intervening protein sequence, to a C-

terminal 113 amino acid activation domain (AD) that can activate transcription when bound to DNA. See, e.g., J. Ma, & M. Ptashne, (1987), Deletion analysis of GALA defines two transcriptional activating segments, Cell 48:847-853: and, J. Ma, & M. Ptashne, (1987), The carboxy-terminal 30 amino acids of GALA are recognized by GAL80, Cell 50:137-142.

- By placing amber codons towards, e.g., the N-terminal DBD of a single polypeptide that contains both the N-terminal DBD of GAL4 and its C-terminal AD, amber suppression by the O-tRNA/O-RS pair can be linked to transcriptional activation by GAL4 (Figure 1, Panel A). GAL4 activated reporter genes can be used to perform both positive and negative selections with the gene (Figure 1, Panel B).
- 10 [0125] The medium used for negative selection can comprise a selecting or screening agent that is converted to a detectable substance by the negative selection marker. In one aspect of the invention, the detectable substance is a toxic substance. A polynucleotide that encodes a negative selection marker can be, e.g., an ura3 gene. For example, the URA3 reporter can be placed under control of a promoter that contains GAL4 DNA binding sites. When the negative selection marker is produced, e.g., by translation of 15 a polynucleotide encoding the GAL4 with selector codons, GAL4 activates transcription of URA3. The negative selection is accomplished on a medium that comprises 5-fluoroorotic acid (5-FOA), which is converted into a detectable substance (e.g., a toxic substance which kills the cell) by the gene product of the ura3 gene. See, e.g., J.D. Boeke, et al., (1984), A 20 positive selection for mutants lacking orotidine-5'-phosphate decarboxylase activity in yeast: 5-fluoroorotic acid resistance, Molecular & General Genetics 197:345-346); M. Vidal, et al., (1996), Genetic characterization of a mammalian protein-protein interaction domain by using a yeast reverse two-hybrid system.[comment], Proc. Natl. Acad. Sci. U. S. A. 93:10321-10326; and, M. Vidal, et al., (1996), Reverse two-hybrid and one-hybrid systems to detect dissociation of protein-protein and DNA-protein interactions.[comment], 25 Proc. Natl. Acad. Sci. U. S. A. 93:10315-10320. See also, Figure 8C.
  - As with the positive selection marker, the negative selection marker can also be any of a variety of molecules. In one embodiment, the positive selection marker and/or the negative selection marker is a polypeptide that fluoresces or catalyzes a luminescent reaction in the presence of a suitable reactant. For example, negative selection markers include, but are not limited to, e.g., luciferase, green fluorescent protein (GFP), YFP, EGFP, RFP, the product of an antibiotic resistant gene (e.g., chloramphenicol acetyltransferase (CAT)), the product of a lacZ gene, transcriptional modulator protein, etc. In one aspect of

the invention, the positive selection marker and/or the negative selection marker is detected by fluorescence-activated cell sorting (FACS) or by luminescence. In another example, the positive selection marker and/or negative selection marker comprise an affinity based screening marker. The same polynucleotide can encode both the positive selection marker and the negative selection marker.

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[0127] Additional levels of selection/screening stringency can also be used in the methods of the invention. The selection or screening stringency can be varied on one or both steps of the method to produce an O-RS. This could include, e.g., varying the amount of response elements in a polynucleotide that encodes the positive and/or negative selection marker, adding a varying amount of an inactive synthetase to one or both of the steps, varying the amount of selection/screening agent that is used, etc. Additional rounds of positive and/or negative selections can also be performed.

[0128] Selecting or screening can also comprise one or more positive or negative selection or screening that includes, e.g., a change in amino acid permeability, a change in translation efficiency, a change in translational fidelity, etc. Typically, the one or more change is based upon a mutation in one or more polynucleotides that comprise or encode components of an orthogonal tRNA-tRNA synthetase pair that are used to produce protein.

[0129] Model enrichment studies can also be used to rapidly select an active synthetase from an excess of inactive synthetases. Positive and/or negative model selection studies can be done. For example, eukaryotic cells that comprise potential active aminoacyl-tRNA synthetases are mixed with a varying fold excess of inactive aminoacyl-tRNA synthetases. A ratio comparison is made between cells grown in a nonselective media and assayed by, e.g., X-GAL overlay, and those grown and able to survive in a selective media (e.g., in the absence of histidine and/or uracil) and assayed by, e.g., an X-GAL assay. For a negative model selection, potential active aminoacyl-tRNA synthetases are mixed with a varying fold excess of inactive aminoacyl-tRNA synthetases and selection is performed with a negative selection substance, e.g., 5-FOA.

[0130] Typically, the library of RSs (e.g., a library of mutant RSs) comprises RSs derived from at least one aminoacyl-tRNA synthetase (RS), e.g., from a non-eukaryotic organism. In one embodiment, the library of RSs is derived from an inactive RS, e.g., where the inactive RS is generated by mutating an active RS, e.g., at the active site in the synthetase, at the editing mechanism site in the synthetase, at different sites by combining

different domains of synthetases, or the like. For example, residues in the active site of the RS are mutated to, e.g., alanine residues. The polynucleotide that encodes the alanine mutated RS is used as a template to mutagenize the alanine residues to all 20 amino acids. The library of mutant RSs is selected/screened to produce the O-RS. In another embodiment, the inactive RS comprises an amino acid binding pocket and one or more amino acids that comprise the binding pocket are substituted with one or more different amino acids. In one example, the substituted amino acids are substituted with alanines. Optionally, the polynucleotide that encodes the alanine mutated RS is used as a template to mutagenize the alanine residues to all 20 amino acids and screened/selected.

of RSs by using various mutagenesis techniques known in the art. For example, the mutant RSs can be generated by site-specific mutations, random point mutations, homologous recombination, DNA shuffling or other recursive mutagenesis methods, chimeric construction or any combination thereof. For example, a library of mutant RSs can be produced from two or more other, e.g., smaller, less diverse "sub-libraries." Once the synthetases are subject to the positive and negative selection/screening strategy, these synthetases can then be subjected to further mutagenesis. For example, a nucleic acid that encodes the O-RS can be isolated; a set of polynucleotides that encode mutated O-RSs (e.g., by random mutagenesis, site-specific mutagenesis, recombination or any combination thereof) can be generated from the nucleic acid; and, these individual steps or a combination of these steps can be repeated until a mutated O-RS is obtained that preferentially aminoacylates the O-tRNA with the unnatural amino acid. In one aspect of the invention, the steps are performed at least two times.

entitled "Methods and compositions for the production of orthogonal tRNA-aminoacyltRNA synthetase pairs." See also, Hamano-Takaku et al., (2000) A mutant Escherichia coli Tyrosyl-tRNA Synthetase Utilizes the Unnatural Amino Acid Azatyrosine More Efficiently than Tyrosine, Journal of Biological Chemistry, 275(51):40324-40328; Kiga et al. (2002), An engineered Escherichia coli tyrosyl-tRNA synthetase for site-specific incorporation of an unnatural amino acid into proteins in eukaryotic translation and its application in a wheat germ cell-free system, PNAS 99(15): 9715-9723; and, Francklyn et al., (2002), Aminoacyl-tRNA synthetases: Versatile players in the changing theater of translation; RNA, 8:1363-1372.

# **ORTHOGONAL tRNAs**

[0133] Eukaryotic cells that include an orthogonal tRNA (O-tRNA) are provided by the invention. The orthogonal tRNA mediates incorporation of an unnatural amino acid into a protein that is encoded by a polynucleotide that comprises a selector codon that is recognized by the O-tRNA, in vivo. In certain embodiments, an O-tRNA of the invention mediates the incorporation of an unnatural amino acid into a protein with, e.g., at least 40%, at least 45%, at least 50%, at least 60%, at least 75%, at least 80%, or even 90% or more as efficiently as tRNA that comprises or is processed in a cell from a polynucleotide sequence as set forth in SEQ ID NO.: 65. See, Table 5, herein.

Example 6 and Table 5, herein). SEQ ID NO.: 65 is a pre-splicing/processing transcript that is optionally processed in the cell, e.g., using the cell's endogenous splicing and processing machinery, and modified to form an active O-tRNA. Typically, a population of such presplicing transcripts form a population of active tRNAs in the cell (the active tRNAs can be in one or more active forms). The invention also includes conservative variations of the O-tRNA and its processed cellular products. For example, conservative variations of O-tRNA include those molecules that function like the O-tRNA of SEQ ID NO.:65 and maintain the tRNA L-shaped structure, e.g., in processed form, but do not have the same sequence (and are other than wild type tRNA molecules). Typically, an O-tRNA of the invention is a recyclable O-tRNA, because the O-tRNA can be reaminoacylated in vivo to again mediate the incorporation of the unnatural amino acid into a protein that is encoded by a polynucleotide in response to a selector codon.

[0135] The transcription of the tRNA in eukaryotes, but not in prokaryotes, is carried out by RNA Polymerase III, which places restrictions on the primary sequence of the tRNA structural genes that can be transcribed in eukaryotic cells. In addition, in eukaryotic cells, tRNAs need to be exported from the nucleus, where they are transcribed, to the cytoplasm, to function in translation. Nucleic acids that encode an O-tRNA of the invention or a complementary polynucleotide thereof are also a feature of the invention. In one aspect of the invention, a nucleic acid that encodes an O-tRNA of the invention includes an internal promoter sequence, e.g., an A box (e.g., TRGCNNAGY) and a B box (e.g., GGTTCGANTCC, SEQ ID NO:88). The O-tRNA of the invention can also be post-transcriptionally modified. For example, post-transcriptional modification of tRNA genes in eukaryotes include removal of the 5'- and 3'- flanking sequences by Rnase P and a 3'-

endonuclease, respectively. The addition of a 3'- CCA sequence is also a post-transcriptional modification of a tRNA gene in eukaryotes.

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[0136] In one embodiment, an O-tRNA is obtained by subjecting to negative selection a population of eukaryotic cells of a first species, where the eukaryotic cells comprise a member of a library of tRNAs. The negative selection eliminates cells that comprise a member of the library of tRNAs that is aminoacylated by an aminoacyl-tRNA synthetase (RS) that is endogenous to the eukaryotic cells. This provides a pool of tRNAs that are orthogonal to the eukaryotic cell of the first species.

[0137] Alternatively, or in combination with others methods described above to incorporate an unnatural amino acid into a polypeptide, a trans-translation system can be used. This system involves a molecule called tmRNA present in *Escherichia coli*. This RNA molecule is structurally related to an alanyl tRNA and is aminoacylated by the alanyl synthetase. The difference between tmRNA and tRNA is that the anticodon loop is replaced with a special large sequence. This sequence allows the ribosome to resume translation on sequences that have stalled using an open reading frame encoded within the tmRNA as template. In the invention, an orthogonal tmRNA can be generated that is preferentially aminoacylated with an orthogonal synthetase and loaded with an unnatural amino acid. By transcribing a gene by the system, the ribosome stalls at a specific site; the unnatural amino acid is introduced at that site, and translation resumes using the sequence encoded within the orthogonal tmRNA.

[0138] Additional methods for producing a recombinant orthogonal tRNAs can be found, e.g., in International patent applications WO 2002/086075, entitled "Methods and compositions for the production of orthogonal tRNA-aminoacyltRNA synthetase pairs." See also, Forster et al., (2003) Programming peptidomimetic synthetases by translating genetic codes designed de novo PNAS 100(11):6353-6357; and, Feng et al., (2003), Expanding tRNA recognition of a tRNA synthetase by a single amino acid change, PNAS 100(10): 5676-5681.

# ORTHOGONAL trna and orthogonal aminoacyl-trna synthetase Pairs

[0139] An orthogonal pair is composed of an O-tRNA, e.g., a suppressor tRNA, a frameshift tRNA, or the like, and an O-RS. The O-tRNA is not acylated by endogenous synthetases and is capable of mediating incorporation of an unnatural amino acid into a

protein that is encoded by a polynucleotide that comprises a selector codon that is recognized by the O-tRNA in vivo. The O-RS recognizes the O-tRNA and preferentially aminoacylates the O-tRNA with an unnatural amino acid in a eukaryotic cell. Methods for producing orthogonal pairs along with orthogonal pairs produced by such methods and compositions of orthogonal pairs for use in eukaryotic cells are included in the invention. The development of multiple orthogonal tRNA/synthetase pairs can allow the simultaneous incorporation of multiple unnatural amino acids using different codons in a eukaryotic cell.

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[0140] An orthogonal O-tRNA/O-RS pair in a eukaryotic cell can be produced by importing a pair, e.g., a nonsense suppressor pair, from a different organism with inefficient cross species aminoacylation. The O-tRNA and O-RS are efficiently expressed and processed in the eukaryotic cell and the O-tRNA is efficiently exported from the nucleus to the cytoplasm. For example, one such pair is the tyrosyl-tRNA synthetase/tRNA<sub>CUA</sub> pair from E. coli (see, e.g., H. M. Goodman, et al., (1968), Nature 217:1019-24; and, D. G. Barker, et al., (1982), FEBS Letters 150:419-23). E. coli tyrosyl-tRNA synthetase efficiently aminoacylates its cognate E. coli tRNA<sub>CUA</sub> when both are expressed in the cytoplasm of S. cerevisiae, but does not aminoacylate S. cerevisiae tRNAs. See, e.g., H. Edwards, & P. Schimmel, (1990), Molecular & Cellular Biology 10:1633-41; and, H. Edwards, et al., (1991), PNAS United States of America 88:1153-6. In addition, E. coli tyrosyl tRNA<sub>CUA</sub> is a poor substrate for S. cerevisiae aminoacyl-tRNA synthetases (see, e.g., V. Trezeguet, et al., (1991), Molecular & Cellular Biology 11:2744-51), but functions efficiently in protein translation in S. cerevisiae. See, e.g., H. Edwards, & P. Schimmel, (1990) Molecular & Cellular Biology 10:1633-41; H. Edwards, et al., (1991), PNAS United States of America 88:1153-6; and, V. Trezeguet, et al., (1991), Molecular & Cellular Biology 11:2744-51. Moreover, E. coli TyrRS does not have an editing mechanism to proofread an unnatural amino acid ligated to the tRNA.

[0141] The O-tRNA and O-RS can be naturally occurring or can be derived by mutation of a naturally occurring tRNA and/or RS, which generates libraries of tRNAs and/or libraries of RSs, from a variety of organism. See the section entitled "Sources and Hosts" herein. In various embodiments, the O-tRNA and O-RS are derived from at least one organism. In another embodiment, the O-tRNA is derived from a naturally occurring or mutated naturally occurring RS from a second organism. In one

embodiment, the first and second non-eukaryotic organisms are the same. Alternatively, the first and second non-eukaryotic organisms can be different.

[0142] See sections herein entitled "Orthogonal aminoacyl-tRNA synthetases" and "O-tRNA" for methods of producing O-RSs and O-tRNAs. *See also*, International patent application WO 2002/086075, entitled "Methods and compositions for the production of orthogonal tRNA-aminoacyltRNA synthetase pairs."

## FIDELITY, EFFICIENCY, AND YIELD

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[0143] Fidelity refers to the accuracy with which a desired molecule, e.g., an unnatural amino acid or amino acid, is incorporated into a growing polypeptide at a desired position. The translational components of the invention incorporate unnatural amino acids, with high fidelity, into proteins in response to a selector codon. For example, using the components of the invention, the efficiency of incorporation of a desired unnatural amino acid into a growing polypeptide chain at a desired position (e.g., in response to a selector codon) is, e.g., greater than 75%, greater than 85%, greater than 95%, or even greater than 99% or more as efficient as compared to unwanted incorporation a specific natural amino acid being incorporated into the growing polypeptide chain the desired position.

[0144] Efficiency can also refer to the degree with which the O-RS aminoacylates the O-tRNA with the unnatural amino acid as compared to a relevant control. O-RSs of the invention can be defined by their efficiency. In certain embodiments of the invention, an O-RS is compared to another O-RS. For example, a O-RS of the invention aminoacylates a O-tRNA with an unnatural amino acid, e.g., at least 40%, at least 50%, at least 60%, at least 75%, at least 80%, at least 90%, at least 95%, or even 99% or more as efficiently as an O-RS having an amino acid sequence, e.g., as set forth in SEQ ID NO.: 86 or 45 )or another specific RS in Table 5) aminoacylates an O-tRNA. In another embodiment, an O-RS of the invention aminoacylates the O-tRNA with the unnatural amino acid at least 10-fold, at least 20-fold, at least 30-fold, etc., more efficiently than the O-RS aminoacylates the O-tRNA with a natural amino acid.

[0145] Using the translational components of the invention, the yield of the polypeptide of interest comprising the unnatural amino acid is, e.g., at least 5%, at least 10%, at least 20%, at least 30%, at least 40%, 50% or more, of that obtained for the naturally occurring polypeptide of interest from a cell in which the polynucleotide lacks the selector codon. In another aspect, the cell produces the polypeptide of interest in the

absence of the unnatural amino acid with a yield that is, e.g., less than 30%, less than 20%, less than 15%, less than 15%, less than 2.5%, etc., of the yield of the polypeptide in the presence of the unnatural amino acid.

## SOURCE AND HOST ORGANISMS

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- 5 [0146] The orthogonal translational components of the invention are typically derived from non-eukaryotic organisms for use in eukaryotic cells or translation systems. For example, the orthogonal O-tRNA can be derived from a non-eukaryotic organism, e.g., a eubacterium, such as Escherichia coli, Thermus thermophilus, Bacillus stearothermphilus, or the like, or an archaebacterium, such as Methanococcus jannaschii, Methanobacterium 10 thermoautotrophicum, Halobacterium such as Haloferax volcanii and Halobacterium species NRC-1, Archaeoglobus fulgidus, Pyrococcus furiosus, Pyrococcus horikoshii, Aeuropyrum pernix, or the like, while the orthogonal O-RS can be derived from a noneukaryotic organism, e.g., a eubacterium, such as Escherichia coli, Thermus thermophilus, Bacillus stearothermphilus, or the like, or an archaebacterium, such as Methanococcus 15 jannaschii, Methanobacterium thermoautotrophicum, Halobacterium such as Haloferax volcanii and Halobacterium species NRC-1, Archaeoglobus fulgidus, Pyrococcus furiosus, Pyrococcus horikoshii, Aeuropyrum pernix, or the like. Alternately, eukaryotic sources can also be used, e.g., plants, algae, protists, fungi, yeasts, animals (e.g., mammals, insects, arthropods, etc.), or the like, e.g., where the components are orthogonal to a cell or translation system of interest, or where they are modified (e.g., mutated) to be orthogonal to 20 the cell or translation system.
  - [0147] The individual components of an O-tRNA/O-RS pair can be derived from the same organism or different organisms. In one embodiment, the O-tRNA/O-RS pair is from the same organism. For example, the O-tRNA/O-RS pair can be derived from a tyrosyltRNA synthetase/tRNA<sub>CUA</sub> pair from *E. coli*. Alternatively, the O-tRNA and the O-RS of the O-tRNA/O-RS pair are optionally from different organisms.
  - [0148] The orthogonal O-tRNA, O-RS or O-tRNA/O-RS pair can be selected or screened and/or used in a eukaryotic cell to produce a polypeptide with an unnatural amino acid. A eukaryotic cell can be from an of a variety of sources, e.g., a plant (e.g., complex plant such as monocots, or dicots), an algae, a protist, a fungus, a yeast (e.g., Saccharomyces cerevisiae), an animal (e.g., a mammal, an insect, an arthropod, etc.), or the

like. Compositions of eukaryotic cells with translational components of the invention are also a feature of the invention.

[0149] The invention also provides for the efficient screening in one species for optional use in that species and/or a second species (optionally, without additional selection/screening). For example, the components of the O-tRNA/O-RS are selected or screened in one species, e.g., an easily manipulated species (such as a yeast cell, etc.) and introduced into a second eukaryotic species, e.g., a plant (e.g., complex plant such as monocots, or dicots), an algae, a protist, a fungus, a yeast, an animal (e.g., a mammal, an insect, an arthropod, etc.), or the like, for use in the in vivo incorporation of an unnatural amino acid in the second species.

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[0150] For example, Saccharomyces cerevisiae (S. cerevisiae) can be chosen as the eukaryotic first species, as it is unicellular, has a rapid generation time, and relatively wellcharacterized genetics. See, e.g., D. Burke, et al., (2000) Methods in Yeast Genetics. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY. Moreover, since the translational machinery of eukaryotes is highly conserved (see, e.g., (1996) Translational Control. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY; Y. Kwok, & J.T. Wong, (1980), Evolutionary relationship between Halobacterium cutirubrum and eukaryotes determined by use of aminoacyl-tRNA synthetases as phylogenetic probes, Canadian Journal of Biochemistry 58:213-218; and, (2001) The Ribosome. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY), aaRSs genes for the incorporation of unnatural amino acids discovered in S. cerevisiae can be introduced into higher eukaryotic organisms and used, in partnership with cognate tRNAs (see, e.g., K. Sakamoto, et al., (2002) Site-specific incorporation of an unnatural amino acid into proteins in mammalian cells, Nucleic Acids Res. 30:4692-4699; and, C. Kohrer, et al., (2001), Import of amber and ochre suppressor tRNAs into mammalian cells: a general approach to site-specific insertion of amino acid analogues into proteins, Proc. Natl. Acad. Sci. U. S. A. 98:14310-14315) to incorporate unnatural amino acids.

[0151] In one example, the method of producing O-tRNA/O-RS in a first species as described herein further includes introducing a nucleic acid that encodes the O-tRNA and a nucleic acid that encodes the O-RS into a eukaryotic cell of a second species (e.g., a mammal, an insect, a fungus, an algae, a plant and the like). In another example, a method of producing an orthogonal aminoacyl-tRNA synthetase (O-RS) that preferentially

aminoacylates an orthogonal tRNA with an unnatural amino acid in a eukaryotic cell includes: (a) subjecting to positive selection, in the presence of an unnatural amino acid, a population of eukaryotic cells of a first species (e.g., yeast and the like). Each of the eukaryotic cells comprise: i) a member of a library of aminoacyl-tRNA synthetases (RSs), ii) an orthogonal tRNA (O-tRNA), iii) a polynucleotide that encodes a positive selection marker, and iv) a polynucleotide that encodes a negative selection marker. The cells that survive the positive selection comprise an active RS that aminoacylates the orthogonal tRNA (O-tRNA) in the presence of an unnatural amino acid. The cells that survive the positive selection are subjected to negative selection in the absence of the unnatural amino acid to eliminate active RSs that aminoacylate the O-tRNA with a natural amino acid. This provides an O-RS that preferentially aminoacylates the O-tRNA with the unnatural amino acid. A nucleic acid that encodes the O-tRNA and a nucleic acid that encodes the O-RS (or the components O-tRNA and/or O-RS) are introduced into a eukaryotic cell of a second species e.g., a mammal, an insect, a fungus, an algae, a plant and/or the like. Typically, the O-tRNA is obtained by subjecting to negative selection a population of eukaryotic cells of a first species, where the eukaryotic cells comprise a member of a library of tRNAs. The negative selection eliminates cells that comprise a member of the library of tRNAs that is aminoacylated by an aminoacyl-tRNA synthetase (RS) that is endogenous to the eukaryotic cells, which provides a pool of tRNAs that are orthogonal to the eukaryotic cell of the first species and the second species.

#### **SELECTOR CODONS**

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[0152] Selector codons of the invention expand the genetic codon framework of the protein biosynthetic machinery. For example, a selector codon includes, e.g., a unique three base codon, a nonsense codon, such as a stop codon, e.g., an amber codon (UAG), an opal codon (UGA), an unnatural codon, at least a four base codon, a rare codon, or the like. A number of selector codons can be introduced into a desired gene, e.g., one or more, two or more, more than three, etc. Once gene can include multiple copies of a given selector codon, or can include multiple different selector codons, or any combination thereof.

[0153] In one embodiment, the methods involve the use of a selector codon that is a stop codon for the incorporation of unnatural amino acids in vivo in a eukaryotic cell. For example, an O-tRNA is produced that recognizes the stop codon, e.g., UAG, and is aminoacylated by an O-RS with a desired unnatural amino acid. This O-tRNA is not

recognized by the naturally occurring host's aminoacyl-tRNA synthetases. Conventional site-directed mutagenesis can be used to introduce the stop codon, e.g., TAG, at the site of interest in a polypeptide of interest. See, e.g., Sayers, J.R., et al. (1988), 5',3' Exonuclease in phosphorothioate-based oligonucleotide-directed mutagenesis. Nucleic Acids Res, 791-802. When the O-RS, O-tRNA and the nucleic acid that encodes the polypeptide of interest are combined in vivo, the unnatural amino acid is incorporated in response to the UAG codon to give a polypeptide containing the unnatural amino acid at the specified position.

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The incorporation of unnatural amino acids in vivo can be done without significant perturbation of the eukaryotic host cell. For example, because the suppression efficiency for the UAG codon depends upon the competition between the O-tRNA, e.g., the amber suppressor tRNA, and a eukaryotic release factor (e.g., eRF) (which binds to a stop codon and initiates release of the growing peptide from the ribosome), the suppression efficiency can be modulated by, e.g., increasing the expression level of O-tRNA, e.g., the suppressor tRNA.

[0155]Selector codons also comprise extended codons, e.g., four or more base codons, such as, four, five, six or more base codons. Examples of four base codons include, e.g., AGGA, CUAG, UAGA, CCCU and the like. Examples of five base codons include, e.g., AGGAC, CCCCU, CCCUC, CUAGA, CUACU, UAGGC and the like. A feature of the invention includes using extended codons based on frameshift suppression. Four or more base codons can insert, e.g., one or multiple unnatural amino acids into the same protein. For example, in the presence of mutated O-tRNAs, e.g., a special frameshift suppressor tRNAs, with anticodon loops, e.g., with at least 8-10 nt anticodon loops, the four or more base codon is read as single amino acid. In other embodiments, the anticodon loops can decode, e.g., at least a four-base codon, at least a five-base codon, or at least a six-base codon or more. Since there are 256 possible four-base codons, multiple unnatural amino acids can be encoded in the same cell using a four or more base codon. See, Anderson et al., (2002) Exploring the Limits of Codon and Anticodon Size, Chemistry and Biology, 9:237-244; Magliery, (2001) Expanding the Genetic Code: Selection of Efficient Suppressors of Four-base Codons and Identification of "Shifty" Four-base Codons with a Library Approach in Escherichia coli, J. Mol. Biol. 307: 755-769.

[0156] For example, four-base codons have been used to incorporate unnatural amino acids into proteins using in vitro biosynthetic methods. See, e.g., Ma et al., (1993)

Biochemistry, 32:7939; and Hohsaka et al., (1999) J. Am. Chem. Soc., 121:34. CGGG and AGGU were used to simultaneously incorporate 2-naphthylalanine and an NBD derivative of lysine into streptavidin in vitro with two chemically acylated frameshift suppressor tRNAs. See, e.g., Hohsaka et al., (1999) J. Am. Chem. Soc., 121:12194. In an in vivo study, Moore et al. examined the ability of tRNALeu derivatives with NCUA anticodons to suppress UAGN codons (N can be U, A, G, or C), and found that the quadruplet UAGA can be decoded by a tRNALeu with a UCUA anticodon with an efficiency of 13 to 26% with little decoding in the 0 or -1 frame. See, Moore et al., (2000) J. Mol. Biol., 298:195. In one embodiment, extended codons based on rare codons or nonsense codons can be used in invention, which can reduce missense readthrough and frameshift suppression at other unwanted sites.

[0157] For a given system, a selector codon can also include one of the natural three base codons, where the endogenous system does not use (or rarely uses) the natural base codon. For example, this includes a system that is lacking a tRNA that recognizes the natural three base codon, and/or a system where the three base codon is a rare codon.

[0158] Selector codons optionally include unnatural base pairs. These unnatural base pairs further expand the existing genetic alphabet. One extra base pair increases the number of triplet codons from 64 to 125. Properties of third base pairs include stable and selective base pairing, efficient enzymatic incorporation into DNA with high fidelity by a polymerase, and the efficient continued primer extension after synthesis of the nascent unnatural base pair. Descriptions of unnatural base pairs which can be adapted for methods and compositions include, e.g., Hirao, et al., (2002) An unnatural base pair for incorporating amino acid analogues into protein, Nature Biotechnology, 20:177-182. Other relevant publications are listed below.

[0159] For in vivo usage, the unnatural nucleoside is membrane permeable and is phosphorylated to form the corresponding triphosphate. In addition, the increased genetic information is stable and not destroyed by cellular enzymes. Previous efforts by Benner and others took advantage of hydrogen bonding patterns that are different from those in canonical Watson-Crick pairs, the most noteworthy example of which is the iso-C:iso-G pair. See, e.g., Switzer et al., (1989) J. Am. Chem. Soc., 111:8322; and Piccirilli et al., (1990) Nature, 343:33; Kool, (2000) Curr. Opin. Chem. Biol., 4:602. These bases in general mispair to some degree with natural bases and cannot be enzymatically replicated.

Kool and co-workers demonstrated that hydrophobic packing interactions between bases can replace hydrogen bonding to drive the formation of base pair. See, Kool, (2000) Curr. Opin. Chem. Biol., 4:602; and Guckian and Kool, (1998) Angew. Chem. Int. Ed. Engl., 36, 2825. In an effort to develop an unnatural base pair satisfying all the above requirements, Schultz, Romesberg and co-workers have systematically synthesized and studied a series of 5 unnatural hydrophobic bases. A PICS:PICS self-pair is found to be more stable than natural base pairs, and can be efficiently incorporated into DNA by Klenow fragment of Escherichia coli DNA polymerase I (KF). See, e.g., McMinn et al., (1999) J. Am. Chem. Soc., 121:11586; and Ogawa et al., (2000) J. Am. Chem. Soc., 122:3274. A 3MN:3MN self-pair can be synthesized by KF with efficiency and selectivity sufficient for biological 10 function. See, e.g., Ogawa et al., (2000) J. Am. Chem. Soc., 122:8803. However, both bases act as a chain terminator for further replication. A mutant DNA polymerase has been recently evolved that can be used to replicate the PICS self pair. In addition, a 7AI self pair can be replicated. See, e.g., Tae et al., (2001) J. Am. Chem. Soc., 123:7439. A novel metallobase pair, Dipic:Py, has also been developed, which forms a stable pair upon binding 15 Cu(II). See, Meggers et al., (2000) J. Am. Chem. Soc., 122:10714. Because extended codons and unnatural codons are intrinsically orthogonal to natural codons, the methods of the invention can take advantage of this property to generate orthogonal tRNAs for them.

[0160] A translational bypassing system can also be used to incorporate an unnatural amino acid in a desired polypeptide. In a translational bypassing system, a large sequence is inserted into a gene but is not translated into protein. The sequence contains a structure that serves as a cue to induce the ribosome to hop over the sequence and resume translation downstream of the insertion.

# **UNNATURAL AMINO ACIDS**

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25 [0161] As used herein, an unnatural amino acid refers to any amino acid, modified amino acid, or amino acid analogue other than selenocysteine and/or pyrrolysine and the following twenty genetically encoded alpha-amino acids: alanine, arginine, asparagine, aspartic acid, cysteine, glutamine, glutamic acid, glycine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, valine.

The generic structure of an alpha-amino acid is illustrated by Formula I:

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[0162] An unnatural amino acid is typically any structure having Formula I wherein the R group is any substituent other than one used in the twenty natural amino acids. See, e.g., *Biochemistry* by L. Stryer, 3<sup>rd</sup> ed. 1988, Freeman and Company, New York, for structures of the twenty natural amino acids. Note that, the unnatural amino acids of the invention can be naturally occurring compounds other than the twenty alpha-amino acids above.

Because the unnatural amino acids of the invention typically differ from the [0163] natural amino acids in side chain, the unnatural amino acids form amide bonds with other amino acids, e.g., natural or unnatural, in the same manner in which they are formed in naturally occurring proteins. However, the unnatural amino acids have side chain groups that distinguish them from the natural amino acids. For example, R in Formula I optionally comprises an alkyl-, aryl-, acyl-, keto-, azido-, hydroxyl-, hydrazine, cyano-, halo-, hydrazide, alkenyl, alkynyl, ether, thiol, seleno-, sulfonyl-, borate, boronate, phospho, phosphono, phosphine, heterocyclic, enone, imine, aldehyde, ester, thioacid, hydroxylamine, amine, and the like, or any combination thereof. Other unnatural amino acids of interest include, but are not limited to, amino acids comprising a photoactivatable cross-linker, spin-labeled amino acids, fluorescent amino acids, metal binding amino acids, metal-containing amino acids, radioactive amino acids, amino acids with novel functional groups, amino acids that covalently or noncovalently interact with other molecules, photocaged and/or photoisomerizable amino acids, biotin or biotin-analogue containing amino acids, keto containing amino acids, amino acids comprising polyethylene glycol or polyether, heavy atom substituted amino acids, chemically cleavable or photocleavable amino acids, amino acids with an elongated side chain as compared to natural amino acids (e.g., polyethers or long chain hydrocarbons, e.g., greater than about 5, greater than about 10 carbons, etc.), carbon-linked sugar-containing amino acids, redox-active amino acids, amino thioacid containing amino acids, and amino acids containing one or more toxic moiety. In some embodiments, the unnatural amino acids have a photoactivatable crosslinker that is used, e.g., to link a protein to a solid support. In one embodiment, the

unnatural amino acids have a saccharide moiety attached to the amino acid side chain (e.g., glycosylated amino acids) and/or other carbohydrate modification.

[0164] In addition to unnatural amino acids that contain novel side chains, unnatural amino acids also optionally comprise modified backbone structures, e.g., as illustrated by the structures of Formula II and III:

$$\prod_{z \leftarrow c \rightarrow \forall H}$$

wherein Z typically comprises OH, NH<sub>2</sub>, SH, NH-R', or S-R'; X and Y, which can be the same or different, typically comprise S or O, and R and R', which are optionally the same or different, are typically selected from the same list of constituents for the R group described above for the unnatural amino acids having Formula I as well as hydrogen. For example, unnatural amino acids of the invention optionally comprise substitutions in the amino or carboxyl group as illustrated by Formulas II and III. Unnatural amino acids of this type include, but are not limited to,  $\alpha$ -hydroxy acids,  $\alpha$ -thioacids  $\alpha$ -aminothiocarboxylates, e.g., with side chains corresponding to the common twenty natural amino acids or unnatural side chains. In addition, substitutions at the  $\alpha$ -carbon optionally include L, D, or  $\alpha$ - $\alpha$ -disubstituted amino acids such as D-glutamate, D-alanine, D-methyl-O-tyrosine, aminobutyric acid, and the like. Other structural alternatives include cyclic amino acids, such as proline analogues as well as 3,4,6,7,8, and 9 membered ring proline analogues,  $\beta$  and  $\gamma$  amino acids such as substituted  $\beta$ -alanine and  $\gamma$ -amino butyric acid.

[0165] For example, many unnatural amino acids are based on natural amino acids, such as tyrosine, glutamine, phenylalanine, and the like. Tyrosine analogs include parasubstituted tyrosines, ortho-substituted tyrosines, and meta substituted tyrosines, where the substituted tyrosine comprises, e.g., a keto group (e.g., an acetyl group), a benzoyl group, an 5 amino group, a hydrazine, an hydroxyamine, a thiol group, a carboxy group, an isopropyl group, a methyl group, a C<sub>6</sub> - C<sub>20</sub> straight chain or branched hydrocarbon, a saturated or unsaturated hydrocarbon, an O-methyl group, a polyether group, a nitro group, an alkynyl group or the like. In addition, multiply substituted aryl rings are also contemplated. Glutamine analogs of the invention include, but are not limited to, α-hydroxy derivatives, γsubstituted derivatives, cyclic derivatives, and amide substituted glutamine derivatives. 10 Example phenylalanine analogs include, but are not limited to, para-substituted phenylalanines, ortho-substituted phenyalanines, and meta-substituted phenylalanines, where the substituent comprises, e.g., a hydroxy group, a methoxy group, a methyl group, an allyl group, an aldehyde, an azido, an iodo, a bromo, a keto group (e.g., an acetyl group), 15 a benzoyl, an alkynyl group, or the like. Specific examples of unnatural amino acids include, but are not limited to, a p-acetyl-L- phenylalanine, a p-propargyloxyphenylalanine, O-methyl-L-tyrosine, an L-3-(2-naphthyl)alanine, a 3-methyl-phenylalanine, an O-4-allyl-Ltyrosine, a 4-propyl-L-tyrosine, a tri-O-acetyl-GlcNAcβ-serine, an L-Dopa, a fluorinated phenylalanine, an isopropyl-L-phenylalanine, a p-azido-L-phenylalanine, a p-acyl-Lphenylalanine, a p-benzoyl-L-phenylalanine, an L-phosphoserine, a phosphonoserine, a 20 phosphonotyrosine, a p-iodo-phenylalanine, a p-bromophenylalanine, a p-amino-Lphenylalanine, and an isopropyl-L-phenylalanine, and the like. Examples of structures of unnatural amino acids are illustrated in Figure 7, Panel B and Figure 11. Additional structures of a variety of unnatural amino acids are provided in, for example, Figures 16, 17, 18, 19, 26, and 29 of WO 2002/085923 entitled "In vivo incorporation of unnatural amino 25 acids." See also, figure 1 structures 2-5 of Kiick et al., (2002) Incorporation of azides into recombinant proteins for chemoselective modification by the Staudinger ligitation, PNAS 99:19-24, for additional methionine analogs.

[0166] In one embodiment, compositions that include an unnatural amino acid (such as p-(propargyloxy)-phenyalanine) are provided. Various compositions comprising p-(propargyloxy)-phenyalanine and, e.g., proteins and/or cells, are also provided. In one aspect, a composition that includes the p-(propargyloxy)-phenyalanine unnatural amino acid further includes an orthogonal tRNA. The unnatural amino acid can be bonded (e.g.,

covalently) to the orthogonal tRNA, e.g., covalently bonded to the orthogonal tRNA though an amino-acyl bond, covalently bonded to a 3'OH or a 2'OH of a terminal ribose sugar of the orthogonal tRNA, etc.

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The chemical moieties via an unnatural amino acids that can be incorporated [0167] into proteins offer a variety of advantages and manipulations of the protein. For example, the unique reactivity of a keto functional group allows selective modification of proteins with any of a number of hydrazine- or hydroxylamine-containing reagents in vitro and in vivo. A heavy atom unnatural amino acid, for example, can be useful for phasing x-ray structure data. The site-specific introduction of heavy atoms using unnatural amino acids also provides selectivity and flexibility in choosing positions for heavy atoms. Photoreactive unnatural amino acids (e.g., amino acids with benzophenone and arylazides (e.g., phenylazide) side chains), for example, allow for efficient in vivo and in vitro photocrosslinking of proteins. Examples of photoreactive unnatural amino acids include, but are not limited to, e.g., p-azido-phenylalanine and p-benzoyl-phenylalanine. The protein with the photoreactive unnatural amino acids can then be crosslinked at will by excitation of the photoreactive group-providing temporal (and/or spatial) control. In one example, the methyl group of an unnatural amino can be substituted with an isotopically labeled, e.g., methyl group, as a probe of local structure and dynamics, e.g., with the use of nuclear magnetic resonance and vibrational spectroscopy. Alkynyl or azido functional groups, for example, allow the selective modification of proteins with molecules through a [3+2] cycloaddition reaction.

# Chemical Synthesis of Unnatural Amino Acids

[0168] Many of the unnatural amino acids provided above are commercially available, e.g., from Sigma (USA) or Aldrich (Milwaukee, WI, USA). Those that are not commercially available are optionally synthesized as provided herein or as provided in various publications or using standard methods known to those of skill in the art. For organic synthesis techniques, see, e.g., Organic Chemistry by Fessendon and Fessendon, (1982, Second Edition, Willard Grant Press, Boston Mass.); Advanced Organic Chemistry by March (Third Edition, 1985, Wiley and Sons, New York); and Advanced Organic Chemistry by Carey and Sundberg (Third Edition, Parts A and B, 1990, Plenum Press, New York). Additional publications describing the synthesis of unnatural amino acids include, e.g., WO 2002/085923 entitled "In vivo incorporation of Unnatural Amino Acids;" Matsoukas et al., (1995) J. Med. Chem., 38, 4660-4669; King, F.E. & Kidd, D.A.A. (1949)

A New Synthesis of Glutamine and of y-Dipeptides of Glutamic Acid from Phthylated Intermediates. J. Chem. Soc., 3315-3319; Friedman, O.M. & Chatterrji, R. (1959) Synthesis of Derivatives of Glutamine as Model Substrates for Anti-Tumor Agents. J. Am. Chem. Soc. 81, 3750-3752; Craig, J.C. et al. (1988) Absolute Configuration of the Enantiomers of 7-Chloro-4 [[4-(diethylamino)-1-methylbutyl]amino]quinoline (Chloroquine). J. Org. Chem. 5 53, 1167-1170; Azoulay, M., Vilmont, M. & Frappier, F. (1991) Glutamine analogues as Potential Antimalarials,. Eur. J. Med. Chem. 26, 201-5; Koskinen, A.M.P. & Rapoport, H. (1989) Synthesis of 4-Substituted Prolines as Conformationally Constrained Amino Acid Analogues. J. Org. Chem. 54, 1859-1866; Christie, B.D. & Rapoport, H. (1985) Synthesis of 10 Optically Pure Pipecolates from L-Asparagine. Application to the Total Synthesis of (+)-Apovincamine through Amino Acid Decarbonylation and Iminium Ion Cyclization. J. Org. Chem. 1989:1859-1866; Barton et al., (1987) Synthesis of Novel a-Amino-Acids and Derivatives Using Radical Chemistry: Synthesis of L- and D-a-Amino-Adipic Acids, L-aaminopimelic Acid and Appropriate Unsaturated Derivatives. Tetrahedron Lett. 43:4297-15 4308; and, Subasinghe et al., (1992) Quisqualic acid analogues: synthesis of betaheterocyclic 2-aminopropanoic acid derivatives and their activity at a novel quisqualatesensitized site. J. Med. Chem. 35:4602-7. See also, patent application entitled "Protein Arrays," attorney docket number P1001US00 filed on December 22, 2002.

[0169] In one aspect of the invention, a method for synthesizing a *p*(propargyloxy)phenyalanine compound is provided. A method comprises, e.g., (a)
suspending N-tert-butoxycarbonyl-tyrosine and K<sub>2</sub>CO<sub>3</sub> in anhydrous DMF; (b) adding
propargyl bromide to the reaction mixture of (a) and alkylating the hydroxyl and the
carboxyl group, resulting in an protected intermediate compound having the structure:

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and (c) mixing the protected intermediate compound with anhydrous HCl in MeOH and deprotecting the amine moiety, thereby synthesizing the *p*-(propargyloxy)phenyalanine compound. In one embodiment, the method further comprises (d) dissolving the p-(propargyloxy)phenylalanine HCl in aqueous NaOH and MeOH and stirring it at room temperature; (e) adjusting the pH of to pH 7; and (f) precipitating the p
(propargyloxy)phenylalanine compound. See e.g., synthesis of propargyloxyphenylalanine in Example 4, herein.

# Cellular uptake of unnatural amino acids

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[0170] Unnatural amino acid uptake by a eukaryotic cell is one issue that is typically considered when designing and selecting unnatural amino acids, e.g., for incorporation into a protein. For example, the high charge density of α-amino acids suggests that these compounds are unlikely to be cell permeable. Natural amino acids are taken up into the eukaryotic cell via a collection of protein-based transport systems. A rapid screen can be done which assesses which unnatural amino acids, if any, are taken up by cells. *See*, e.g., the toxicity assays in, e.g., the application entitled "Protein Arrays," attorney docket number P1001US00 filed on December 22, 2002; and Liu, D.R. & Schultz, P.G. (1999) *Progress toward the evolution of an organism with an expanded genetic code*. PNAS United States 96:4780-4785. Although uptake is easily analyzed with various assays, an alternative to designing unnatural amino acids that are amenable to cellular uptake pathways is to provide biosynthetic pathways to create amino acids *in vivo*.

## Biosynthesis of Unnatural Amino Acids

Many biosynthetic pathways already exist in cells for the production of [0171] amino acids and other compounds. While a biosynthetic method for a particular unnatural amino acid may not exist in nature, e.g., in a eukaryotic cell, the invention provides such methods. For example, biosynthetic pathways for unnatural amino acids are optionally generated in host cell by adding new enzymes or modifying existing host cell pathways. Additional new enzymes are optionally naturally occurring enzymes or artificially evolved enzymes. For example, the biosynthesis of p-aminophenylalanine (as presented in an example in WO 2002/085923 entitled "In vivo incorporation of unnatural amino acids") relies on the addition of a combination of known enzymes from other organisms. The genes for these enzymes can be introduced into a eukaryotic cell by transforming the cell with a plasmid comprising the genes. The genes, when expressed in the cell, provide an enzymatic pathway to synthesize the desired compound. Examples of the types of enzymes that are optionally added are provided in the examples below. Additional enzymes sequences are found, e.g., in Genbank. Artificially evolved enzymes are also optionally added into a cell in the same manner. In this manner, the cellular machinery and resources of a cell are manipulated to produce unnatural amino acids.

[0172] A variety of methods are available for producing novel enzymes for use in biosynthetic pathways or for evolution of existing pathways. For example, recursive recombination, e.g., as developed by Maxygen, Inc. (available on the world wide web at

www.maxygen.com), is optionally used to develop novel enzymes and pathways. See, e.g., Stemmer (1994), Rapid evolution of a protein in vitro by DNA shuffling, Nature 370(4):389-391; and, Stemmer, (1994), DNA shuffling by random fragmentation and reassembly: In vitro recombination for molecular evolution, Proc. Natl. Acad. Sci. USA., 91:10747-10751. Similarly DesignPath™, developed by Genencor (available on the world wide web at genencor.com) is optionally used for metabolic pathway engineering, e.g., to engineer a pathway to create O-methyl-L-tyrosine in a cell. This technology reconstructs existing pathways in host organisms using a combination of new genes, e.g., identified through functional genomics, and molecular evolution and design. Diversa Corporation (available on the world wide web at diversa.com) also provides technology for rapidly screening libraries of genes and gene pathways, e.g., to create new pathways.

[0173] Typically, the unnatural amino acid produced with an engineered biosynthetic pathway of the invention is produced in a concentration sufficient for efficient protein biosynthesis, e.g., a natural cellular amount, but not to such a degree as to affect the concentration of the other amino acids or exhaust cellular resources. Typical concentrations produced *in vivo* in this manner are about 10 mM to about 0.05 mM. Once a cell is transformed with a plasmid comprising the genes used to produce enzymes desired for a specific pathway and an unnatural amino acid is generated, *in vivo* selections are optionally used to further optimize the production of the unnatural amino acid for both ribosomal protein synthesis and cell growth.

#### POLYPEPTIDES WITH UNNATURAL AMINO ACIDS

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[0174] Proteins or polypeptides of interest with at least one unnatural amino acid are a feature of the invention. The invention also includes polypeptides or proteins with at least one unnatural amino acid produced using the compositions and methods of the invention. An excipient (e.g., a pharmaceutically acceptable excipient) can also be present with the protein.

By producing proteins or polypeptides of interest with at least one unnatural amino acid in eukaryotic cells, proteins or polypeptides will typically include eukaryotic posttranslational modifications. In certain embodiments, a protein includes at least one unnatural amino acid and at least one post-translational modification that is made in vivo by a eukaryotic cell, where the post-translational modification is not made by a prokaryotic cell. For example, the post-translation modification includes, e.g., acetylation, acylation,

lipid-modification, palmitoylation, palmitate addition, phosphorylation, glycolipid-linkage modification, glycosylation, and the like. In one aspect, the post-translational modification includes attachment of an oligosaccharide (e.g., (GlcNAc-Man)<sub>2</sub>-Man-GlcNAc-GlcNAc)) to an asparagine by a GlcNAc-asparagine linkage. See also, Table 7, which lists some examples of N-linked oligosaccharides of eukaryotic proteins (additional residues can also be present, which are not shown). In another aspect, the post-translational modification includes attachment of an oligosaccharide (e.g., Gal-GalNAc, Gal-GlcNAc, etc.) to a serine or threonine by a GalNAc-serine or GalNAc-threonine linkage, or a GlcNAc-serine or a GlcNAc-threonine linkage.

TABLE 7: EXAMPLES OF OLIGOSACCHARIDES THROUGH GICNAC-LINKAGE

Type	Base Structure
High-mannose	Manα1-6 Manα1-3 Manα1-6 Manα1-3 Manα1-4GlcNAcβ1-4GlcNAcβ1-Asn
Hybrid	Manα1-6 Manβ1-4GlcNAcβ1-4GlcNAcβ1-Asn GlcNAcβ1-2 — Manα1-3
Complex	GlcNAc $\beta$ 1-2 — Man $\alpha$ 1-6 — Man $\beta$ 1-4GlcNAc $\beta$ 1-4GlcNAc $\beta$ 1-Asn GlcNAc $\beta$ 1-2 — Man $\alpha$ 1-3
Xylose	Manα1-6 Manβ1-4GlcNAcβ1-4GlcNAcβ1-Asn Xylβ1-2

In yet another aspect, the post-translation modification includes proteolytic processing of precursors (e.g., calcitonin precursor, calcitonin gene-related peptide precursor, preproparathyroid hormone, preproinsulin, proinsulin, prepro-opiomelanocortin, pro-opiomelanocortin and the like), assembly into a multisubunit protein or macromolecular assembly, translation to another site in the cell (e.g., to organelles, such as the endoplasmic reticulum, the golgi apparatus, the nucleus, lysosomes, peroxisomes, mitochondria, chloroplasts, vacuoles, etc., or through the secretory pathway). In certain embodiments, the protein comprises a secretion or localization sequence, an epitope tag, a FLAG tag, a polyhistidine tag, a GST fusion, or the like.

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[0177] One advantage of an unnatural amino acid is that it presents additional chemical moieties that can be used to add additional molecules. These modifications can be made in vivo in a eukaryotic cell, or in vitro. Thus, in certain embodiments, the post-translational modification is through the unnatural amino acid. For example, the post-translational modification can be through a nucleophilic-electrophilic reaction. Most reactions currently used for the selective modification of proteins involve covalent bond formation between nucleophilic and electrophilic reaction partners, e.g. the reaction of  $\alpha$ -haloketones with histidine or cysteine side chains. Selectivity in these cases is determined

by the number and accessibility of the nucleophilic residues in the protein. In proteins of the invention, other more selective reactions can be used, such as the reaction of an unnatural keto-amino acid with hydrazides or aminooxy compounds, in vitro and in vivo. See, e.g., Cornish, et al., (1996) Am. Chem. Soc., 118:8150-8151; Mahal, et al., (1997) Science, 276:1125-1128; Wang, et al., (2001) Science 292:498-500; Chin, et al., (2002) Am. Chem. Soc. 124:9026-9027; Chin, et al., (2002) Proc. Natl. Acad. Sci., 99:11020-11024; Wang, et al., (2003) Proc. Natl. Acad. Sci., 100:56-61; Zhang, et al., (2003) Biochemistry, 42:6735-6746; and, Chin, et al., (2003) Science, in press. This allows the selective labeling of virtually any protein with a host of reagents including fluorophores, crosslinking agents, saccharide derivatives and cytotoxic molecules. See also, patent application USSN 10/686,944 entitled "Glycoprotein synthesis" filed October 15, 2003. Post-translational modifications, e.g., through an azido amino acid, can also made through the Staudinger ligation (e.g., with triarylphosphine reagents). See, e.g., Kiick et al., (2002) Incorporation of azides into recombinant proteins for chemoselective modification by the Staudinger ligation, PNAS 99:19-24.

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This invention provides another highly efficient method for the selective [0178] modification of proteins, which involves the genetic incorporation of unnatural amino acids, e.g., containing an azide or alkynyl moiety (see, e.g., 2 and 1 of Figure 11), into proteins in response to a selector codon. These amino acid side chains can then be modified by, e.g., a 20 Huisgen [3+2] cycloaddition reaction (see, e.g., Padwa, A. in Comprehensive Organic Synthesis, Vol. 4, (1991) Ed. Trost, B. M., Pergamon, Oxford, p. 1069-1109; and, Huisgen, R. in 1,3-Dipolar Cycloaddition Chemistry, (1984) Ed. Padwa, A., Wiley, New York, p. 1-176) with, e.g., alkynyl or azide derivatives, respectively. See, e.g., Figure 16. Because this method involves a cycloaddition rather than a nucleophilic substitution, proteins can be 25 modified with extremely high selectivity. This reaction can be carried out at room temperature in aqueous conditions with excellent regions electivity (1,4 > 1,5) by the addition of catalytic amounts of Cu(I) salts to the reaction mixture. See, e.g., Tornoe, et al., (2002) Org. Chem. 67:3057-3064; and, Rostovtsev, et al., (2002) Angew. Chem. Int. Ed. 41:2596-2599. Another method that can be used is the ligand exchange on a bisarsenic compound with a tetracysteine motif, see, e.g., Griffin, et al., (1998) Science 281:269-272. 30

[0179] A molecule that can be added to a protein of the invention through a [3+2] cycloaddition includes virtually any molecule with an azido or alkynyl derivative. See, e.g., Example 3 and 5, herein. Such molecules include, but are not limited to, dyes,

fluorophores, crosslinking agents, saccharide derivatives, polymers (e.g., derivatives of polyethylene glycol), photocrosslinkers, cytotoxic compounds, affinity labels, derivatives of biotin, resins, beads, a second protein or polypeptide (or more), polynucleotide(s) (e.g., DNA, RNA, etc.), metal chelators, cofactors, fatty acids, carbohydrates, and the like. *See*, *e.g.*, **Figures 13A**, and Example 3 and 5, herein. These molecules can be added to an unnatural amino acid with an alkynyl group, e.g., p-propargyloxyphenylalanine, or azido group, e.g., p-azido-phenylalanine, respectively. For example, see **Figure 13B** and **Figure 17A**.

[0180] In another aspect, the invention provides compositions including such molecules and methods of producing these molecules, e.g., azido dyes (such as shown in chemical structure 4 and chemical structure 6), an alkynyl polyethylene glycol (e.g., as shown in chemical structure 7), where n is an integer between, e.g., 50 and 10,000, 75 and 5,000, 100 and 2,000, 100 and 1,000, etc. In embodiment of the invention, the alkynyl polyethylene glycol has a molecular weight of, e.g., about 5,000 to about 100,000 Da, about 20,000 to about 50, 000 Da, about 20,000 to about 10,000 Da), etc.

[0181]Various compositions comprising these compounds, e.g., with proteins and cells, are also provided. In one aspect of the invention, a protein comprising an azido dye (e.g., of chemical structure 4 or chemical structure 6), further includes at least one unnatural amino acid (e.g., an alkynyl amino acid), where the azido dye is attached to the unnatural amino acid through a [3+2] cycloaddition.

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[0182]In one embodiment, a protein comprises the alkynyl polyethylene glycol of chemical structure 7. In another embodiment, the composition further includes at least one unnatural amino acid (e.g., an azido amino acid), wherein the alkynyl polyethylene glycol is attached to an unnatural amino acid through a [3+2] cycloaddition.

Methods for synthesizing azido dyes are also provided. For example, one [0183] such method comprises: (a) providing a dye compound comprising a sulfonyl halide moiety; (b) warming the dye compound to room temperature in the presence of 3-azidopropylamine and triethylamine and coupling an amine moiety of the 3-azidopropylamine to the halide position of the dye compound, thereby synthesizing the azido dye. In one example embodiment, the dye compound comprises dansyl chloride, and the azido dye comprises the 15 composition of chemical structure 4. In one aspect, the method further comprises purifying the azido dye from the reaction mixture. See, e.g., Example 5, herein.

In another example, a method for synthesizing an azido dye comprises (a) [0184] providing an amine-containing dye compound; (b) combining the amine-containing dye compound with a carbodiimide and 4-(3-azidopropylcarbamoyl)-butyric acid in a suitable solvent, and coupling a carbonyl group of the acid to the amine moiety of the dye compound, thereby synthesizing the azido dye. In one embodiment, the carbodiimine comprises 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDCI). In one aspect, the amine-containing dye comprises fluoresceinamine, and the suitable solvent comprises pyridine. For example, the amine-containing dye optionally comprises fluoresceinamine and the azido dye optionally comprises the composition of chemical structure 6. In one embodiment, the method further comprises (c) precipitating the azido dye; (d) washing the precipitate with HCl; (e) dissolving the washed precipitate in EtOAc; and (f) precipitating the azido dye in hexanes. See, e.g., Example 5, herein.

Methods for synthesizing a propargyl amide polyethylene glycol are also [0185] provided. For example, the method comprises reacting propargylamine with polyethylene glycol (PEG)-hydroxysuccinimide ester in an organic solvent (e.g., CH<sub>2</sub>Cl<sub>2</sub>) at room

temperature, resulting in the propargyl amide polyethylene glycol of chemical structure 7. In one embodiment, the method further comprises precipitating the propargylamide polyethylene glycol using ethyl acetate. In one aspect, the method further includes recrystallizing the propargylamide polyethylene glycol in methanol; and drying the product under a vacuum. *See, e.g.*, Example 5, herein.

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[0186] A eukaryotic cell of the invention provides the ability to synthesize proteins that comprise unnatural amino acids in large useful quantities. In one aspect, the composition optionally includes, e.g., at least 10 micrograms, at least 50 micrograms, at least 75 micrograms, at least 100 micrograms, at least 200 micrograms, at least 250 micrograms, at least 500 micrograms, at least 1 milligram, at least 10 milligrams or more of the protein that comprises an unnatural amino acid, or an amount that can be achieved with in vivo protein production methods (details on recombinant protein production and purification are provided herein). In another aspect, the protein is optionally present in the composition at a concentration of, e.g., at least 10 micrograms of protein per liter, at least 50 micrograms of protein per liter, at least 75 micrograms of protein per liter, at least 100 micrograms of protein per liter, at least 200 micrograms of protein per liter, at least 250 micrograms of protein per liter, at least 500 micrograms of protein per liter, at least 1 milligram of protein per liter, or at least 10 milligrams of protein per liter or more, in, e.g., a cell lysate, a buffer, a pharmaceutical buffer, or other liquid suspension (e.g., in a volume of, e.g., anywhere from about 1 nl to about 100 L). The production of large quantities (e.g., greater that that typically possible with other methods, e.g., in vitro translation) of a protein in a eukaryotic cell including at least one unnatural amino acid is a feature of the invention.

[0187] The incorporation of an unnatural amino acid can be done to, e.g., tailor changes in protein structure and/or function, e.g., to change size, acidity, nucleophilicity, hydrogen bonding, hydrophobicity, accessibility of protease target sites, target to a moiety (e.g., for a protein array), etc. Proteins that include an unnatural amino acid can have enhanced or even entirely new catalytic or physical properties. For example, the following properties are optionally modified by inclusion of an unnatural amino acid into a protein: toxicity, biodistribution, structural properties, spectroscopic properties, chemical and/or photochemical properties, catalytic ability, half-life (e.g., serum half-life), ability to react with other molecules, e.g., covalently or noncovalently, and the like. The compositions including proteins that include at least one unnatural amino acid are useful for, e.g., novel therapeutics, diagnostics, catalytic enzymes, industrial enzymes, binding proteins (e.g.,

antibodies), and e.g., the study of protein structure and function. See, e.g., Dougherty, (2000) Unnatural Amino Acids as Probes of Protein Structure and Function, Current Opinion in Chemical Biology, 4:645-652.

In one aspect of the invention, a composition includes at least one protein with at least one, e.g., at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, or at least ten or more unnatural amino acids. The unnatural amino acids can be the same or different, e.g., there can be 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 or more different sites in the protein that comprise 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 or more different unnatural amino acids. In another aspect, a composition includes a protein with at least one, but fewer than all, of a particular amino acid present in the protein is substituted with the unnatural amino acids. For a given protein with more than one unnatural amino acids, the unnatural amino acids can be identical or different (e.g., the protein can include two or more different types of unnatural amino acids, or can include two of the same unnatural amino acid. For a given protein with more than two unnatural amino acids, the unnatural amino acids can be the same, different or a combination of a multiple unnatural amino acid of the same kind with at least one different unnatural amino acid.

Essentially any protein (or portion thereof) that includes an unnatural amino acid (and any corresponding coding nucleic acid, e.g., which includes one or more selector codons) can be produced using the compositions and methods herein. No attempt is made to identify the hundreds of thousands of known proteins, any of which can be modified to include one or more unnatural amino acid, e.g., by tailoring any available mutation methods to include one or more appropriate selector codon in a relevant translation system.

Common sequence repositories for known proteins include GenBank EMBL, DDBJ and the NCBI. Other repositories can easily be identified by searching the internet.

[0190] Typically, the proteins are, e.g., at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, at least 95%, or at least 99% or more identical to any available protein (e.g., a therapeutic protein, a diagnostic protein, an industrial enzyme, or portion thereof, and the like), and they comprise one or more unnatural amino acid. Examples of therapeutic, diagnostic, and other proteins that can be modified to comprise one or more unnatural amino acids include, but are not limited to, e.g., Alpha-1 antitrypsin, Angiostatin, Antihemolytic factor, antibodies (further details on antibodies are found below), Apolipoprotein, Apoprotein, Atrial natriuretic factor, Atrial natriuretic polypeptide, Atrial

peptides, C-X-C chemokines (e.g., T39765, NAP-2, ENA-78, Gro-a, Gro-b, Gro-c, IP-10, GCP-2, NAP-4, SDF-1, PF4, MIG), Calcitonin, CC chemokines (e.g., Monocyte chemoattractant protein-1, Monocyte chemoattractant protein-2, Monocyte chemoattractant protein-3, Monocyte inflammatory protein-1 alpha, Monocyte inflammatory protein-1 beta, RANTES, I309, R83915, R91733, HCC1, T58847, D31065, T64262), CD40 ligand, C-kit 5 Ligand, Collagen, Colony stimulating factor (CSF), Complement factor 5a, Complement inhibitor, Complement receptor 1, cytokines, (e.g., epithelial Neutrophil Activating Peptide-78, GROa/MGSA, GROB, GROY, MIP-1\alpha, MIP-1\delta, MCP-1), Epidermal Growth Factor (EGF), Erythropoietin ("EPO", representing a preferred target for modification by the 10 incorporation of one or more unnatural amino acid), Exfoliating toxins A and B, Factor IX, Factor VII, Factor VIII, Factor X, Fibroblast Growth Factor (FGF), Fibringen, Fibronectin, G-CSF, GM-CSF, Glucocerebrosidase, Gonadotropin, growth factors, Hedgehog proteins (e.g., Sonic, Indian, Desert), Hemoglobin, Hepatocyte Growth Factor (HGF), Hirudin, Human serum albumin, Insulin, Insulin-like Growth Factor (IGF), interferons (e.g., IFN-α, IFN-β, IFN-γ), interleukins (e.g., IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, 15 IL-11, IL-12, etc.), Keratinocyte Growth Factor (KGF), Lactoferrin, leukemia inhibitory factor, Luciferase, Neurturin, Neutrophil inhibitory factor (NIF), oncostatin M, Osteogenic protein, Parathyroid hormone, PD-ECSF, PDGF, peptide hormones (e.g., Human Growth Hormone), Pleiotropin, Protein A, Protein G, Pyrogenic exotoxins A, B, and C, Relaxin, 20 Renin, SCF, Soluble complement receptor I, Soluble I-CAM 1, Soluble interleukin receptors (IL-1, 2, 3, 4, 5, 6, 7, 9, 10, 11, 12, 13, 14, 15), Soluble TNF receptor, Somatomedin, Somatostatin, Somatotropin, Streptokinase, Superantigens, i.e., Staphylococcal enterotoxins (SEA, SEB, SEC1, SEC2, SEC3, SED, SEE), Superoxide dismutase (SOD), Toxic shock syndrome toxin (TSST-1), Thymosin alpha 1, Tissue plasminogen activator, Tumor necrosis factor beta (TNF beta), Tumor necrosis factor 25 receptor (TNFR), Tumor necrosis factor-alpha (TNF alpha), Vascular Endothelial Growth Factor (VEGEF), Urokinase, and many others.

[0191] One class of proteins that can be made using the compositions and methods for in vivo incorporation of unnatural amino acids described herein includes transcriptional modulators or portions thereof. Example transcriptional modulators include genes and transcriptional modulator proteins that modulate cell growth, differentiation, regulation, or the like. Transcriptional modulators are found in prokaryotes, viruses, and eukaryotes, including fungi, plants, yeasts, insects, and animals, including mammals, providing a wide

range of therapeutic targets. It will be appreciated that expression and transcriptional activators regulate transcription by many mechanisms, e.g., by binding to receptors, stimulating a signal transduction cascade, regulating expression of transcription factors, binding to promoters and enhancers, binding to proteins that bind to promoters and enhancers, unwinding DNA, splicing pre-mRNA, polyadenylating RNA, and degrading RNA. For example, compositions of GALA protein or portion thereof in a eukaryotic cell are also a feature of the invention. Typically, the GALA protein or portion thereof comprises at least one unnatural amino acid. See also the section herein entitled "Orthogonal aminoacyl-tRNA synthetases."

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[0192] One class of proteins of the invention (e.g., proteins with one or more unnatural amino acids) include expression activators such as cytokines, inflammatory molecules, growth factors, their receptors, and oncogene products, e.g., interleukins (e.g., IL-1, IL-2, IL-8, etc.), interferons, FGF, IGF-I, IGF-II, FGF, PDGF, TNF, TGF-α, TGF-β, EGF, KGF, SCF/c-Kit, CD40L/CD40, VLA-4/VCAM-1, ICAM-1/LFA-1, and
 hyalurin/CD44; signal transduction molecules and corresponding oncogene products, e.g., Mos, Ras, Raf, and Met; and transcriptional activators and suppressors, e.g., p53, Tat, Fos, Myc, Jun, Myb, Rel, and steroid hormone receptors such as those for estrogen, progesterone, testosterone, aldosterone, the LDL receptor ligand and corticosterone.

[0193] Enzymes (e.g., industrial enzymes), or portions thereof with at least one unnatural amino acid, are also provided by the invention. Examples of enzymes include, but are not limited to, e.g., amidases, amino acid racemases, acylases, dehalogenases, dioxygenases, diarylpropane peroxidases, epimerases, epoxide hydrolases, esterases, isomerases, kinases, glucose isomerases, glycosidases, glycosyl transferases, haloperoxidases, monooxygenases (e.g., p450s), lipases, lignin peroxidases, nitrile hydratases, nitrilases, proteases, phosphatases, subtilisins, transaminase, and nucleases.

[0194] Many of these proteins are commercially available (*See*, e.g., the Sigma BioSciences 2002 catalogue and price list), and the corresponding protein sequences and genes and, typically, many variants thereof, are well-known (*see*, e.g., Genbank). Any of them can be modified by the insertion of one or more unnatural amino acid according to the invention, e.g., to alter the protein with respect to one or more therapeutic, diagnostic or enzymatic properties of interest. Examples of therapeutically relevant properties include serum half-life, shelf half-life, stability, immunogenicity, therapeutic activity, detectability

(e.g., by the inclusion of reporter groups (e.g., labels or label binding sites) in the unnatural amino acids), reduction of  $LD_{50}$  or other side effects, ability to enter the body through the gastric tract (e.g., oral availability), or the like. Examples of diagnostic properties include shelf half-life, stability, diagnostic activity, detectability, or the like. Examples of relevant enzymatic properties include shelf half-life, stability, enzymatic activity, production capability, or the like.

[0195] A variety of other proteins can also be modified to include one or more unnatural amino acid of the invention. For example, the invention can include substituting one or more natural amino acids in one or more vaccine proteins with an unnatural amino acid, e.g., in proteins from infectious fungi, e.g., Aspergillus, Candida species; bacteria, particularly *E. coli*, which serves a model for pathogenic bacteria, as well as medically important bacteria such as Staphylococci (e.g., aureus), or Streptococci (e.g., pneumoniae); protozoa such as sporozoa (e.g., Plasmodia), rhizopods (e.g., Entamoeba) and flagellates (Trypanosoma, Leishmania, Trichomonas, Giardia, etc.); viruses such as (+) RNA viruses (examples include Poxviruses e.g., vaccinia; Picornaviruses, e.g. polio; Togaviruses, e.g., rubella; Flaviviruses, e.g., HCV; and Coronaviruses), (-) RNA viruses (e.g., Rhabdoviruses, e.g., VSV; Paramyxovimses, e.g., RSV; Orthomyxovimses, e.g., influenza; Bunyaviruses; and Arenaviruses), dsDNA viruses (Reoviruses, for example), RNA to DNA viruses, i.e., Retroviruses, e.g., HIV and HTLV, and certain DNA to RNA viruses such as Hepatitis B.

[0196] Agriculturally related proteins such as insect resistance proteins (e.g., the Cry proteins), starch and lipid production enzymes, plant and insect toxins, toxin-resistance proteins, Mycotoxin detoxification proteins, plant growth enzymes (e.g., Ribulose 1,5-Bisphosphate Carboxylase/Oxygenase, "RUBISCO"), lipoxygenase (LOX), and Phosphoenolpyruvate (PEP) carboxylase are also suitable targets for unnatural amino acid modification.

[0197] The invention also provides methods for producing in a eukaryotic cell at least one protein comprising at least one unnatural amino acid (and proteins produced by such methods). For example, a method includes: growing, in an appropriate medium, a eukaryotic cell that comprises a nucleic acid that comprises at least one selector codon and encodes the protein. The eukaryotic cell also comprises: an orthogonal tRNA (O-tRNA) that functions in the cell and recognizes the selector codon; and an orthogonal aminoacyl

tRNA synthetase (O-RS) that preferentially aminoacylates the O-tRNA with the unnatural amino acid, and the medium comprises an unnatural amino acid.

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In one embodiment, the method further includes incorporating into the protein the unnatural amino acid, where the unnatural amino acid comprises a first reactive group; and contacting the protein with a molecule (e.g., a dye, a polymer, e.g., a derivative of polyethylene glycol, a photocrosslinker, a cytotoxic compound, an affinity label, a derivative of biotin, a resin, a second protein or polypeptide, a metal chelator, a cofactor, a fatty acid, a carbohydrate, a polynucleotide (e.g., DNA, RNA, etc.), and the like) that comprises a second reactive group. The first reactive group reacts with the second reactive group to attach the molecule to the unnatural amino acid through a [3+2] cycloaddition. In one embodiment, the first reactive group is an alkynyl or azido moiety and the second reactive group is an azido or alkynyl moiety. For example, the first reactive group is the alkynyl moiety (e.g., in unnatural amino acid p-propargyloxyphenylalanine) and the second reactive group is the azido moiety (e.g., in the unnatural amino acid p-azido-L-phenylalanine) and the second reactive group is the alkynyl moiety.

[0199] In one embodiment, the O-RS aminoacylates the O-tRNA with the unnatural amino acid at least 50% as efficiently as does an O-RS having an amino acid sequence, e.g., as set forth in SEQ ID NO.: 86 or 45. In another embodiment, the O-tRNA comprises, is processed from, or is encoded by SEQ ID NO.: 65 or 64, or a complementary polynucleotide sequence thereof. In yet another embodiment, the O-RS comprises an amino acid set forth in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63) and/or 86.

[0200] The encoded protein can comprise, e.g., a therapeutic protein, a diagnostic protein, an industrial enzyme, or portion thereof. Optionally, the protein that is produced by the method is further modified through the unnatural amino acid. For example, the protein produced by the method is optionally modified by at least one post-translational modification in vivo.

[0201] Methods of producing a screening or selecting transcriptional modulator protein are also provided (and screening or selecting transcriptional modulator proteins produced by such methods). For example, a method includes: selecting a first polynucleotide sequence, where the polynucleotide sequence encodes a nucleic acid binding

domain; and mutating the first polynucleotide sequence to include at least one selector codon. This provides a screening or selecting polynucleotide sequence. The method also includes: selecting a second polynucleotide sequence, where the second polynucleotide sequence encodes a transcriptional activation domain; providing a construct that comprises the screening or selecting polynucleotide sequence operably linked to the second polynucleotide sequence; and, introducing the construct, an unnatural amino acid, an orthogonal tRNA synthetase (O-RS) and an orthogonal tRNA (O-tRNA) into a cell. With these components, the O-RS preferentially aminoacylates the O-tRNA with the unnatural amino acid and the O-tRNA recognizes the selector codon and incorporates the unnatural amino acid into the nucleic acid binding domain, in response to the selector codon in the screening or selecting polynucleotide sequence, thereby providing the screening or selecting transcriptional modulator protein.

[0202] In certain embodiments, the protein or polypeptide of interest (or portion thereof) in the methods and/or compositions of the invention is encoded by a nucleic acid. Typically, the nucleic acid comprises at least one selector codon, at least two selector codons, at least three selector codons, at least four selector codons, at least five selector codons, at least six selector codons, at least seven selector codons, at least eight selector codons, at least nine selector codons, ten or more selector codons.

[0203] Genes coding for proteins or polypeptides of interest can be mutagenized using methods well-known to one of skill in the art and described herein under "Mutagenesis and Other Molecular Biology Techniques" to include, e.g., one or more selector codon for the incorporation of an unnatural amino acid. For example, a nucleic acid for a protein of interest is mutagenized to include one or more selector codon, providing for the insertion of the one or more unnatural amino acids. The invention includes any such variant, e.g., mutant, versions of any protein, e.g., including at least one unnatural amino acid. Similarly, the invention also includes corresponding nucleic acids, i.e., any nucleic acid with one or more selector codon that encodes one or more unnatural amino acid.

[0204] In one example embodiment, the invention provides compositions (& compositions produced by the methods of the invention) that include a Thr44, Arg110 TAG mutant of GAL4, where the GAL4 protein includes at least one unnatural amino acid. In another embodiment, the invention provides compositions that include a Trp33 TAG mutant

of human Superoxide dimutase (hSOD), where the hSOD protein includes at least one unnatural amino.

Purifying recombinant proteins comprising unnatural amino acids Proteins of the invention, e.g., proteins comprising unnatural amino acids, [0205] 5 antibodies to proteins comprising unnatural amino acids, etc., can be purified, either partially or substantially to homogeneity, according to standard procedures known to and used by those of skill in the art. Accordingly, polypeptides of the invention can be recovered and purified by any of a number of methods well known in the art, including, e.g., ammonium sulfate or ethanol precipitation, acid or base extraction, column 10 chromatography, affinity column chromatography, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, hydroxylapatite chromatography, lectin chromatography, gel electrophoresis and the like. Protein refolding steps can be used, as desired, in making correctly folded mature proteins. High performance liquid chromatography (HPLC), 15 affinity chromatography or other suitable methods can be employed in final purification steps where high purity is desired. In one embodiment, antibodies made against unnatural amino acids (or proteins comprising unnatural amino acids) are used as purification reagents, e.g., for affinity-based purification of proteins comprising one or more unnatural amino acid(s). Once purified, partially or to homogeneity, as desired, the polypeptides are 20 optionally used e.g., as assay components, therapeutic reagents or as immunogens for antibody production.

[0206] In addition to other references noted herein, a variety of purification/protein folding methods are well known in the art, including, e.g., those set forth in R. Scopes, Protein Purification, Springer-Verlag, N.Y. (1982); Deutscher, Methods in Enzymology Vol. 182: Guide to Protein Purification, Academic Press, Inc. N.Y. (1990); Sandana (1997) Bioseparation of Proteins, Academic Press, Inc.; Bollag et al. (1996) Protein Methods, 2nd Edition Wiley-Liss, NY; Walker (1996) The Protein Protocols Handbook Humana Press, NJ, Harris and Angal (1990) Protein Purification Applications: A Practical Approach IRL Press at Oxford, Oxford, England; Harris and Angal Protein Purification Methods: A Practical Approach IRL Press at Oxford, Oxford, England; Scopes (1993) Protein Purification: Principles and Practice 3rd Edition Springer Verlag, NY; Janson and Ryden (1998) Protein Purification: Principles, High Resolution Methods and Applications, Second

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Edition Wiley-VCH, NY; and Walker (1998) <u>Protein Protocols on CD-ROM</u> Humana Press, NJ; and the references cited therein.

[0207] One advantage of producing a protein or polypeptide of interest with an unnatural amino acid in a eukaryotic cell is that typically the proteins or polypeptides will be folded in their native conformations. However, in certain embodiments of the invention, those of skill in the art will recognize that, after synthesis, expression and/or purification, proteins can possess a conformation different from the desired conformations of the relevant polypeptides. In one aspect of the invention, the expressed protein is optionally denatured and then renatured. This is accomplished, e.g., by adding a chaperonin to the protein or polypeptide of interest, and/or by solubilizing the proteins in a chaotropic agent such as guanidine HCl, etc.

[0208] In general, it is occasionally desirable to denature and reduce expressed polypeptides and then to cause the polypeptides to re-fold into the preferred conformation. For example, guanidine, urea, DTT, DTE, and/or a chaperonin can be added to a translation product of interest. Methods of reducing, denaturing and renaturing proteins are well known to those of skill in the art (see, the references above, and Debinski, et al. (1993) <u>J. Biol. Chem.</u>, 268: 14065-14070; Kreitman and Pastan (1993) <u>Bioconjug. Chem.</u>,4: 581-585; and Buchner, et al., (1992) <u>Anal. Biochem.</u>, 205: 263-270). Debinski, et al., for example, describe the denaturation and reduction of inclusion body proteins in guanidine-DTE. The proteins can be refolded in a redox buffer containing, e.g., oxidized glutathione and L-arginine. Refolding reagents can be flowed or otherwise moved into contact with the one or more polypeptide or other expression product, or vice-versa.

## **ANTIBODIES**

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[0209] In one aspect, the invention provides antibodies to molecules of the invention, e.g., synthetases, tRNA, and proteins comprising unnatural amino acids. Antibodies to molecules of the invention are useful as purification reagents, e.g., for purifying the molecules of the invention. In addition, the antibodies can be used as indicator reagents to indicate the presence of a synthetase, a tRNA, or protein comprising an unnatural amino acid, e.g., to track the presence or location (e.g., in vivo or in situ) of the molecule.

[0210] An antibody of the invention can be a protein comprising one or more polypeptides substantially or partially encoded by immunoglobulin genes or fragments of

immunoglobulin genes. The recognized immunoglobulin genes include the kappa, lambda, alpha, gamma, delta, epsilon and mu constant region genes, as well as myriad immunoglobulin variable region genes. Light chains are classified as either kappa or lambda. Heavy chains are classified as gamma, mu, alpha, delta, or epsilon, which in turn define the immunoglobulin classes, IgG, IgM, IgA, IgD and IgE, respectively. A typical immunoglobulin (e.g., antibody) structural unit comprises a tetramer. Each tetramer is composed of two identical pairs of polypeptide chains, each pair having one "light" (about 25 kD) and one "heavy" chain (about 50-70 kD). The N-terminus of each chain defines a variable region of about 100 to 110 or more amino acids primarily responsible for antigen recognition. The terms variable light chain (VL) and variable heavy chain (V<sub>H</sub>) refer to these light and heavy chains, respectively.

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[0211] Antibodies exist as intact immunoglobulins or as a number of wellcharacterized fragments produced by digestion with various peptidases. Thus, for example, pepsin digests an antibody below the disulfide linkages in the hinge region to produce F(ab')<sub>2</sub>, a dimer of Fab which itself is a light chain joined to V<sub>H</sub>-C<sub>H</sub>1 by a disulfide bond. The F(ab')<sub>2</sub> may be reduced under mild conditions to break the disulfide linkage in the hinge region thereby converting the F(ab')<sub>2</sub>dimer into an Fab' monomer. The Fab' monomer is essentially an Fab with part of the hinge region (see, Fundamental Immunology, 4th addition, W.E. Paul, ed., Raven Press, N.Y. (1999), for a more detailed description of other antibody fragments). While various antibody fragments are defined in terms of the digestion of an intact antibody, one of skill will appreciate that such Fab' fragments, etc. may be synthesized de novo either chemically or by utilizing recombinant DNA methodology. Thus, the term antibody, as used herein, also optionally includes antibody fragments either produced by the modification of whole antibodies or synthesized de novo using recombinant DNA methodologies. Antibodies include single chain antibodies, including single chain Fv (sFv or scFv) antibodies in which a variable heavy and a variable light chain are joined together (directly or through a peptide linker) to form a continuous polypeptide. Antibodies of the invention can be, e.g., polyclonal, monoclonal, chimeric, humanized, single chain, Fab fragments, fragments produced by an Fab expression library, or the like.

[0212] In general, antibodies of the invention are valuable, both as general reagents and as therapeutic reagents in a variety of molecular biological or pharmaceutical processes. Methods of producing polyclonal and monoclonal antibodies are available, and can be

applied to making the antibodies of the invention. A number of basic texts describe standard antibody production processes, including, e.g., Borrebaeck (ed) (1995) Antibody Engineering, 2<sup>nd</sup> Edition Freeman and Company, NY (Borrebaeck); McCafferty et al. (1996) Antibody Engineering, A Practical Approach IRL at Oxford Press, Oxford, England (McCafferty), and Paul (1995) Antibody Engineering Protocols Humana Press, Towata, NJ (Paul); Paul (ed.), (1999) Fundamental Immunology, Fifth edition Raven Press, N.Y.; Coligan (1991) Current Protocols in Immunology Wiley/Greene, NY; Harlow and Lane (1989) Antibodies: A Laboratory Manual Cold Spring Harbor Press, NY; Stites et al. (eds.) Basic and Clinical Immunology (4th ed.) Lange Medical Publications, Los Altos, CA, and references cited therein; Goding (1986) Monoclonal Antibodies: Principles and Practice (2d ed.) Academic Press, New York, NY; and Kohler and Milstein (1975) Nature 256: 495-497.

[0213] A variety of recombinant techniques for antibody preparation which do not rely on, e.g., injection of an antigen into an animal have been developed and can be used in the context of the present invention. For example, it is possible to generate and select libraries of recombinant antibodies in phage or similar vectors. See, e.g., Winter et al. (1994) Making Antibodies by Phage Display Technology Annu. Rev. Immunol. 12:433-55 and the references cited therein for a review. See also, Griffiths and Duncan (1998) Strategies for selection of antibodies by phage display Curr Opin Biotechnol 9: 102-8; Hoogenboom et al. (1998) Antibody phage display technology and its applications Immunotechnology 4: 1-20; Gram et al. (1992) in vitro selection and affinity maturation of antibodies from a naïve combinatorial immunoglobulin library PNAS 89:3576-3580; Huse et al. (1989) Science 246: 1275-1281; and Ward, et al. (1989) Nature 341: 544-546.

[0214] In one embodiment, antibody libraries can include repertoires of V genes (e.g., harvested from populations of lymphocytes or assembled in vitro) which are cloned for display of associated heavy and light chain variable domains on the surface of filamentous bacteriophage. Phage are selected by binding to an antigen. Soluble antibodies are expressed from phage infected bacteria and the antibody can be improved, e.g., via mutagenesis. See e.g., Balint and Larrick (1993) Antibody Engineering by Parsimonious Mutagenesis Gene 137:109-118; Stemmer et al. (1993) Selection of an Active Single Chain Fv Antibody From a Protein Linker Library Prepared by Enzymatic Inverse PCR Biotechniques 14(2):256-65; Crameri et al. (1996) Construction and evolution of antibody-phage libraries by DNA shuffling Nature Medicine 2:100-103; and Crameri and Stemmer

(1995) Combinatorial multiple cassette mutagenesis creates all the permutations of mutant and wildtype cassettes BioTechniques 18:194-195.

[0215] Kits for cloning and expression of recombinant antibody phage systems are also known and available, e.g., the "recombinant phage antibody system, mouse ScFv module," from Amersham-Pharmacia Biotechnology (Uppsala, Sweden). Bacteriophage antibody libraries have also been produced for making high affinity human antibodies by chain shuffling (See, e.g., Marks et al. (1992) By- Passing Immunization: Building High Affinity Human Antibodies by Chain Shuffling Biotechniques 10:779-782. It will also be recognized that antibodies can be prepared by any of a number of commercial services (e.g., Bethyl Laboratories (Montgomery, TX), Anawa (Switzerland), Eurogentec (Belgium and in the US in Philadelphia, PA, etc.) and many others.

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[0216] In certain embodiments, it is useful to "humanize" antibodies of the invention, e.g., where the antibodies are to be administered therapeutically. The use of humanized antibodies tends to reduce the incidence of unwanted immune responses against the therapeutic antibodies (e.g., when the patient is a human). The antibody references above describe humanization strategies. In addition to humanized antibodies, human antibodies are also a feature of the invention. Human antibodies consist of characteristically human immunoglobulin sequences. Human antibodies can be produced in using a wide variety of methods (see, e.g., Larrick et al., U.S. Pat. No. 5,001,065, for a review). A general approach for producing human antibodies by trioma technology is described by Ostberg et al. (1983), Hybridoma 2: 361-367, Ostberg, U.S. Pat. No. 4,634,664, and Engelman et al., U.S. Pat. No. 4,634,666.

[0217] A variety of methods of using antibodies in the purification and detection of proteins are known and can be applied to detecting and purifying proteins comprising unnatural amino acids as noted herein. In general, antibodies are useful reagents for ELISA, western blotting, immunochemistry, affinity chromatograpy methods, SPR, and many other methods. The references noted above provide details on how to perform ELISA assays, western blots, surface plasmon resonance (SPR) and the like.

[0218] In one aspect of the invention, antibodies of the invention themselves include unnatural amino acids, providing the antibodies with properties of interest (e.g., improved half-life, stability, toxicity, or the like). See also, the section herein entitled "Polypeptides with unnatural amino acids." Antibodies account for nearly 50% of all compounds

currently in clinical trials (Wittrup, (1999) *Phage on display* <u>Tibtech</u> 17: 423-424 and antibodies are used ubiquitously as diagnostic reagents. Accordingly, the ability to modify antibodies with unnatural amino acids provides an important tool for modifying these valuable reagents.

[0219] For example, there are many applications of MAbs to the field of diagnostics. Assays range from simple spot tests to more involved methods such as the radio-labeled NR-LU-10 MAb from DuPont Merck Co. used for tumor imaging (Rusch et al. (1993) NR-LU-10 monoclonal antibody scanning. A helpful new adjunct to computed tomography in evaluating non-small-cell lung cancer. J Thorac Cardiovasc Surg 106: 200-4). As noted, MAbs are central reagents for ELISA, western blotting, immunochemistry, affinity chromatograpy methods and the like. Any such diagnostic antibody can be modified to include one or more unnatural amino acid, altering, e.g., the specificity or avidity of the Ab for a target, or altering one or more detectable property, e.g., by including a detectable label (e.g., spectrographic, fluorescent, luminescent, etc.) in the unnatural

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amino acid.

One class of valuable antibody reagents are therapeutic Abs. For example, [0220] antibodies can be tumor-specific MAbs that arrest tumor growth by targeting tumor cells for destruction by antibody-dependent cell-mediated cytotoxicity (ADCC) or complementmediated lysis (CML) (these general types of Abs are sometimes referred to as "magic bullets"). One example is Rituxan, an anti-CD20 MAb for the treatment of Non-Hodgkins 20 lymphoma (Scott (1998) Rituximab: a new therapeutic monoclonal antibody for non-Hodgkin's lymphoma Cancer Pract 6: 195-7). A second example relates to antibodies which interfere with a critical component of tumor growth. Herceptin is an anti-HER-2 monoclonal antibody for treatment of metastatic breast cancer, and provides an example of 25 an antibody with this mechanism of action (Baselga et al. (1998) Recombinant humanized anti-HER2 antibody (Herceptin) enhances the antitumor activity of paclitaxel and doxorubicin against HER2/neu overexpressing human breast cancer xenografts [published erratum appears in Cancer Res (1999) 59(8):2020], Cancer Res 58: 2825-31). A third example relates to antibodies for delivery of cytotoxic compounds (toxins, radionuclides, etc.) directly to a tumor or other site of interest. For example, one application Mab is CYT-30 356, a 90Y-linked antibody that targets radiation directly to prostate tumor cells (Deb et al. (1996) Treatment of hormone-refractory prostate cancer with 90Y-CYT-356 monoclonal antibody Clin Cancer Res 2: 1289-97. A fourth application is antibody-directed enzyme

prodrug therapy, where an enzyme co-localized to a tumor activates a systemicallyadministered pro-drug in the tumor vicinity. For example, an anti-Ep-CAM1 antibody linked to carboxypeptidase A is being developed for treatment of colorectal cancer (Wolfe et al. (1999) Antibody-directed enzyme prodrug therapy with the T268G mutant of human carboxypeptidase A1: in vitro and in vivo studies with prodrugs of methotrexate and the 5 thymidylate synthase inhibitors GW1031 and GW1843 Bioconjug Chem 10: 38-48). Other Abs (e.g., antagonists) are designed to specifically inhibit normal cellular functions for therapeutic benefit. An example is Orthoclone OKT3, an anti-CD3 MAb offered by Johnson and Johnson for reducing acute organ transplant rejection (Strate et al. (1990) Orthoclone OKT3 as first-line therapy in acute renal allograft rejection Transplant Proc 22: 10 219-20. Another class of antibody products are agonists. These Mabs are designed to specifically enhance normal cellular functions for therapeutic benefit. For example, Mabbased agonists of acetylcholine receptors for neurotherapy are under development (Xie et al. (1997) Direct demonstration of MuSK involvement in acetylcholine receptor clustering through identification of agonist ScFv Nat. Biotechnol. 15: 768-71. Any of these antibodies 15 can be modified to include one or more unnatural amino acid to enhance one or more therapeutic property (specificity, avidity, serum-half-life, etc.).

[0221] Another class of antibody products provide novel functions. The main antibodies in this group are catalytic antibodies such as Ig sequences that have been engineered to mimic the catalytic abilities of enzymes (Wentworth and Janda (1998) Catalytic antibodies Curr Opin Chem Biol 2: 138-44. For example, an interesting application involves using the catalytic antibody mAb-15A10 to hydrolyze cocaine in vivo for addiction therapy (Mets et al. (1998) A catalytic antibody against cocaine prevents cocaine's reinforcing and toxic effects in rats Proc Natl Acad Sci U S A 95: 10176-81). Catalytic antibodies can also be modified to include one or more unnatural amino acid to improve one or more property of interest.

### Defining Polypeptides by Immunoreactivity

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[0222] Because the polypeptides of the invention provide a variety of new polypeptide sequences (e.g., comprising unnatural amino acids in the case of proteins synthesized in the translation systems herein, or, e.g., in the case of the novel synthetases herein, novel sequences of standard amino acids), the polypeptides also provide new structural features which can be recognized, e.g., in immunological assays. The generation of antibodies or antibodies which specifically bind the polypeptides of the invention, as well

as the polypeptides which are bound by such antibodies or antisera, are a feature of the invention.

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[0223] For example, the invention includes synthetase proteins that specifically bind to or that are specifically immunoreactive with an antibody or antisera generated against an immunogen comprising an amino acid sequence selected from one or more of (SEQ ID NO: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86). To eliminate cross-reactivity with other homologues, the antibody or antisera is subtracted with available control synthetase homologues, such as the wild-type *E. coli* tyrosyl synthetase (TyrRS) (e.g., SEQ ID NO::2).

In one typical format, the immunoassay uses a polyclonal antiserum which was raised against one or more polypeptide comprising one or more of the sequences corresponding to one or more of SEQ ID NO: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, or a substantial subsequence thereof (i.e., at least about 30% of the full length sequence provided). The set of potential polypeptide immunogens derived from SEQ ID NO: 36-63 and 86 are collectively referred to below as "the immunogenic polypeptides." The resulting antisera is optionally selected to have low cross-reactivity against the control synthetase homologues and any such cross-reactivity is removed, e.g., by immunoabsorbtion, with one or more control synthetase homologues, prior to use of the polyclonal antiserum in the immunoassay.

[0225] In order to produce antisera for use in an immunoassay, one or more of the immunogenic polypeptides is produced and purified as described herein. For example, recombinant protein can be produced in a recombinant cell. An inbred strain of mice (used in this assay because results are more reproducible due to the virtual genetic identity of the mice) is immunized with the immunogenic protein(s) in combination with a standard adjuvant, such as Freund's adjuvant, and a standard mouse immunization protocol (see, e.g., Harlow and Lane (1988) Antibodies, A Laboratory Manual, Cold Spring Harbor Publications, New York, for a standard description of antibody generation, immunoassay formats and conditions that can be used to determine specific immunoreactivity. Additional references and discussion of antibodies is also found herein and can be applied here to make antibodies that define/detect polypeptides by immunoreactivity). Alternatively, one or more synthetic or recombinant polypeptide derived from the sequences disclosed herein is conjugated to a carrier protein and used as an immunogen.

[0226] Polyclonal sera are collected and titered against the immunogenic polypeptide in an immunoassay, for example, a solid phase immunoassay with one or more of the immunogenic proteins immobilized on a solid support. Polyclonal antisera with a titer of  $10^6$  or greater are selected, pooled and subtracted with the control synthetase polypeptides to produce subtracted pooled titered polyclonal antisera.

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[0227] The subtracted pooled titered polyclonal antisera are tested for cross reactivity against the control homologues in a comparative immunoassay. In this comparative assay, discriminatory binding conditions are determined for the subtracted titered polyclonal antisera which result in at least about a 5-10 fold higher signal to noise ratio for binding of the titered polyclonal antisera to the immunogenic synthetase as compared to binding to a control synthetase homologue. That is, the stringency of the binding/washing reaction(s) is/are adjusted by the addition of non-specific competitors such as albumin or non-fat dry milk, and/or by adjusting salt conditions, temperature, and/or the like. These binding/washing conditions are used in subsequent assays for determining whether a test polypeptide (a polypeptide being compared to the immunogenic polypeptides and/ or the control polypeptides) is specifically bound by the pooled subtracted polyclonal antisera. In particular, test polypeptides which show at least a 2-5x higher signal to noise ratio than the control synthetase homologue under discriminatory binding conditions, and at least about a ½ signal to noise ratio as compared to the immunogenic polypeptide(s), shares substantial structural similarity with the immunogenic polypeptide as compared to known synthetases, and is, therefore a polypeptide of the invention.

In another example, immunoassays in the competitive binding format are used for detection of a test polypeptide. For example, as noted, cross-reacting antibodies are removed from the pooled antisera mixture by immunoabsorbtion with the control polypeptides. The immunogenic polypeptide(s) are then immobilized to a solid support which is exposed to the subtracted pooled antisera. Test proteins are added to the assay to compete for binding to the pooled subtracted antisera. The ability of the test protein(s) to compete for binding to the pooled subtracted antisera as compared to the immobilized protein(s) is compared to the ability of the immunogenic polypeptide(s) added to the assay to compete for binding (the immunogenic polypeptides compete effectively with the immobilized immunogenic polypeptides for binding to the pooled antisera). The percent cross-reactivity for the test proteins is calculated, using standard calculations.

[0229] In a parallel assay, the ability of the control proteins to compete for binding to the pooled subtracted antisera is optionally determined as compared to the ability of the immunogenic polypeptide(s) to compete for binding to the antisera. Again, the percent cross-reactivity for the control polypeptides is calculated, using standard calculations.

Where the percent cross-reactivity is at least 5-10x as high for the test polypeptides as compared to the control polypeptides and or where the binding of the test polypeptides is approximately in the range of the binding of the immunogenic polypeptides, the test polypeptides are said to specifically bind the pooled subtracted antisera.

[0230] In general, the immunoabsorbed and pooled antisera can be used in a competitive binding immunoassay as described herein to compare any test polypeptide to the immunogenic and/ or control polypeptide(s). In order to make this comparison, the immunogenic, test and control polypeptides are each assayed at a wide range of concentrations and the amount of each polypeptide required to inhibit 50% of the binding of the subtracted antisera to, e.g., an immobilized control, test or immunogenic protein is determined using standard techniques. If the amount of the test polypeptide required for binding in the competitive assay is less than twice the amount of the immunogenic polypeptide that is required, then the test polypeptide is said to specifically bind to an antibody generated to the immunogenic protein, provided the amount is at least about 5-10x as high as for the control polypeptide.

optionally fully immunosorbed with the immunogenic polypeptide(s) (rather than the control polypeptides) until little or no binding of the resulting immunogenic polypeptide subtracted pooled antisera to the immunogenic polypeptide(s) used in the immunosorbtion is detectable. This fully immunosorbed antisera is then tested for reactivity with the test polypeptide. If little or no reactivity is observed (i.e., no more than 2x the signal to noise ratio observed for binding of the fully immunosorbed antisera to the immunogenic polypeptide), then the test polypeptide is specifically bound by the antisera elicited by the immunogenic protein.

#### PHARMACEUTICAL COMPOSITIONS

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[0232] The polypeptides or proteins of the invention (e.g., synthetases, proteins comprising one or more unnatural amino acid, etc.) are optionally employed for therapeutic uses, e.g., in combination with a suitable pharmaceutical carrier. Such compositions, e.g.,

comprise a therapeutically effective amount of the compound, and a pharmaceutically acceptable carrier or excipient. Such a carrier or excipient includes, but is not limited to, saline, buffered saline, dextrose, water, glycerol, ethanol, and/or combinations thereof. The formulation is made to suit the mode of administration. In general, methods of administering proteins are well known in the art and can be applied to administration of the polypeptides of the invention.

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[0233] Therapeutic compositions comprising one or more polypeptide of the invention are optionally tested in one or more appropriate in vitro and/or in vivo animal models of disease, to confirm efficacy, tissue metabolism, and to estimate dosages, according to methods well known in the art. In particular, dosages can be initially determined by activity, stability or other suitable measures of unnatural herein to natural amino acid homologues (e.g., comparison of an EPO modified to include one or more unnatural amino acids to a natural amino acid EPO), i.e., in a relevant assay.

[0234] Administration is by any of the routes normally used for introducing a molecule into ultimate contact with blood or tissue cells. The unnatural amino acid polypeptides of the invention are administered in any suitable manner, optionally with one or more pharmaceutically acceptable carriers. Suitable methods of administering such polypeptides in the context of the present invention to a patient are available, and, although more than one route can be used to administer a particular composition, a particular route can often provide a more immediate and more effective action or reaction than another route.

[0235] Pharmaceutically acceptable carriers are determined in part by the particular composition being administered, as well as by the particular method used to administer the composition. Accordingly, there is a wide variety of suitable formulations of pharmaceutical compositions of the present invention.

[0236] Polypeptide compositions can be administered by a number of routes including, but not limited to: oral, intravenous, intraperitoneal, intramuscular, transdermal, subcutaneous, topical, sublingual, or rectal means. Unnatural amino acid polypeptide compositions can also be administered via liposomes. Such administration routes and appropriate formulations are generally known to those of skill in the art.

[0237] The unnatural amino acid polypeptide, alone or in combination with other suitable components, can also be made into aerosol formulations (i.e., they can be

"nebulized") to be administered via inhalation. Aerosol formulations can be placed into pressurized acceptable propellants, such as dichlorodifluoromethane, propane, nitrogen, and the like.

[0238] Formulations suitable for parenteral administration, such as, for example, by intraarticular (in the joints), intravenous, intramuscular, intradermal, intraperitoneal, and subcutaneous routes, include aqueous and non-aqueous, isotonic sterile injection solutions, which can contain antioxidants, buffers, bacteriostats, and solutes that render the formulation isotonic with the blood of the intended recipient, and aqueous and non-aqueous sterile suspensions that can include suspending agents, solubilizers, thickening agents, stabilizers, and preservatives. The formulations of packaged nucleic acid can be presented in unit-dose or multi-dose sealed containers, such as ampules and vials.

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[0239] Parenteral administration and intravenous administration are preferred methods of administration. In particular, the routes of administration already in use for natural amino acid homologue therapeutics (e.g., those typically used for EPO, GCSF, GMCSF, IFNs, interleukins, antibodies, and/or any other pharmaceutically delivered protein), along with formulations in current use, provide preferred routes of administration and formulation for the proteins that include unnatural amino acids of the invention (e.g., pegylated variants of current thereputic proteins, etc.).

[0240] The dose administered to a patient, in the context of the present invention, is sufficient to effect a beneficial therapeutic response in the patient over time, or, e.g., to inhibit infection by a pathogen, or other appropriate activity, depending on the application. The dose is determined by the efficacy of a particular composition/formulation, and the activity, stability or serum half-life of the unnatural amino acid polypeptide employed and the condition of the patient, as well as the body weight or surface area of the patient to be treated. The size of the dose is also determined by the existence, nature, and extent of any adverse side-effects that accompany the administration of a particular composition/ formulation, or the like in a particular patient.

[0241] In determining the effective amount of the composition/formulation to be administered in the treatment or prophylaxis of disease (e.g., cancers, inherited diseases, diabetes, AIDS, or the like), the physician evaluates circulating plasma levels, formulation toxicities, progression of the disease, and/or where relevant, the production of anti-unnatural amino acid polypeptide antibodies.

[0242] The dose administered, e.g., to a 70 kilogram patient, is typically in the range equivalent to dosages of currently-used therapeutic proteins, adjusted for the altered activity or serum half-life of the relevant composition. The compositions/formulations of this invention can supplement treatment conditions by any known conventional therapy, including antibody administration, vaccine administration, administration of cytotoxic agents, natural amino acid polypeptides, nucleic acids, nucleotide analogues, biologic response modifiers, and the like.

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[0243] For administration, formulations of the present invention are administered at a rate determined by the LD-50 of the relevant formulation, and/or observation of any side-effects of the unnatural amino acids at various concentrations, e.g., as applied to the mass and overall health of the patient. Administration can be accomplished via single or divided doses.

[0244] If a patient undergoing infusion of a formulation develops fevers, chills, or muscle aches, he/she receives the appropriate dose of aspirin, ibuprofen, acetaminophen or other pain/fever controlling drug. Patients who experience reactions to the infusion such as fever, muscle aches, and chills are premedicated 30 minutes prior to the future infusions with either aspirin, acetaminophen, or, e.g., diphenhydramine. Meperidine is used for more severe chills and muscle aches that do not quickly respond to antipyretics and antihistamines. Treatment is slowed or discontinued depending upon the severity of the reaction.

#### NUCLEIC ACID AND POLYPEPTIDE SEQUENCE AND VARIANTS

[0245] As described above and below, the invention provides for nucleic acid polynucleotide sequences and polypeptide amino acid sequences, e.g., O-tRNAs and O-RSs, and, e.g., compositions and methods comprising said sequences. Examples of said sequences, e.g., O-tRNAs and O-RSs are disclosed herein (see, Table 5, e.g., SEQ ID NO. 3-65, 86, and other than SEQ ID NO.: 1 and 2). However, one of skill in the art will appreciate that the invention is not limited to those sequences disclosed herein, e.g., the Examples and Table 5. One of skill will appreciate that the invention also provides many related and even unrelated sequences with the functions described herein, e.g., encoding an O-tRNA or an O-RS.

[0246] The invention also provides polypeptides (O-RSs) and polynucleotides, e.g., O-tRNA, polynucleotides that encode O-RSs or portions thereof (e.g., the active site of the

synthetase), oligonucleotides used to construct aminoacyl-tRNA synthetase mutants, etc. For example, a polypeptide of the invention includes a polypeptide that comprises an amino acid sequence as shown in any one of SEQ ID NO.: 36-63 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, a polypeptide that comprises an amino acid sequence encoded by a polynucleotide sequence as shown in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), and a polypeptide that is specifically immunoreactive with an antibody specific for a polypeptide that comprises an amino acid sequence as shown in any one of SEQ ID NO.: 36-63, and/or 86, or a polypeptide that comprises an amino acid sequence encoded by a polynucleotide sequence as shown in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35).

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[0247] Also included among the polypeptides of the invention are polypeptides that comprise an amino acid sequence that is at least 90% identical to that of a naturally occurring tyrosyl aminoacyl-tRNA synthetase (TyrRS) (e.g., SEQ ID NO.:2) and comprises two or more amino acids of groups A-E. For example, group A includes valine, isoleucine, leucine, glycine, serine, alanine, or threonine at a position corresponding to Tyr37 of E. coli TyrRS; group B includes aspartate at a position corresponding to Asn126 of E. coli TyrRS; group C includes threonine, serine, arginine, asparagine or glycine at a position corresponding to Asp182 of E. coli TyrRS; group D includes methionine, alanine, valine, or tyrosine at a position corresponding to Phe183 of E. coli TyrRS; and, group E includes serine, methionine, valine, cysteine, threonine, or alanine at a position corresponding to Leu 186 of E. coli TyrRS. Any subset of combinations of these groups are a feature of the invention. For example, in one embodiment, the O-RS has two or more amino acids selected from valine, isoleucine, leucine, or threonine occurs at a position corresponding to Tyr37 of E. coli TyrRS; threonine, serine, arginine, or glycine at a position corresponding to Asp182 of E. coli TyrRS; methionine, or tyrosine at a position corresponding to Phe183 of E. coli TyrRS; and, serine, or alanine at a position corresponding to Leu186 of E. coli TyrRS. In another embodiment, the O-RS includes two more more amino acids selected from glycine, serine, or alanine at a position corresponding to Tyr37 of E. coli TyrRS, aspartate at a position corresponding to Asn126 of E. coli TyrRS, asparagine at a position corresponding to Asp182 of E. coli TyrRS, alanine, or valine, at a position corresponding to Phe183 of E. coli TyrRS, and/or methionine, valine, cysteine, or threonine, at a position corresponding to Leu186 of E. coli TyrRS. Similarly, polypeptides of the invention also include a polypeptide that comprises at least 20 contiguous amino acids of SEQ ID NO.: 3663 (e.g., 36-47, 48-63, or any other subset of 36-63), and/or 86, and two or more amino acid substitutions as indicated above in groups A-E. *See also*, Table 4, Table 6, and/or Table 8, herein. An amino acid sequence comprising a conservative variation of any of the above polypeptides is also included as a polypeptide of the invention.

5 [0248] In one embodiment, a composition includes a polypeptide of the invention and an excipient (e.g., buffer, water, pharmaceutically acceptable excipient, etc.). The invention also provides an antibody or antisera specifically immunoreactive with a polypeptide of the invention.

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[0249] Polynucleotides are also provided in the invention. Polynucleotides of the invention include those that encode proteins or polypeptides of interest of the invention, or that include one or more selector codon, or both. For example, polynucleotides of the invention include, e.g., a polynucleotide comprising a nucleotide sequence as set forth in any one of SEQ ID NO.: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), 64-85; a polynucleotide that is complementary to or that encodes a polynucleotide sequence thereof; and/or a polynucleotide encoding a polypeptide that comprises an amino acid sequence as set forth in any one of SEQ ID NO.: 36-63, and/or 86, or a conservative variation thereof. A polynucleotide of the invention also includes a polynucleotide that encodes a polypeptide of the invention. Similarly, a nucleic acid that hybridizes to a polynucleotide indicated above under highly stringent conditions over substantially the entire length of the nucleic acid is a polynucleotide of the invention.

[0250] A polynucleotide of the invention also includes a polynucleotide that encodes a polypeptide that comprises an amino acid sequence that is at least 90% identical to that of a naturally occurring tyrosyl aminoacyl-tRNA synthetase (TyrRS) (e.g., SEQ ID NO.: 2) and comprises two or more mutations as indicated above in groups A-E (above). A polynucleotide that is that is at least 70%, (or at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 99% or more) identical to a polynucleotide indicated above and/or a polynucleotide comprising a conservative variation of any of the polynucleotides indicated above are also included among the polynucleotides of the invention. See also, Table 4, Table 6, and/or Table 8, herein.

[0251] In certain embodiments, a vector (e.g., a plasmid, a cosmid, a phage, a virus, etc.) comprises a polynucleotide of the invention. In one embodiment, the vector is an expression vector. In another embodiment, the expression vector includes a promoter

operably linked to one or more of the polynucleotides of the invention. In another embodiment, a cell comprises a vector that includes a polynucleotide of the invention.

[0252] One of skill will also appreciate that many variants of the disclosed sequences are included in the invention. For example, conservative variations of the disclosed sequences that yield a functionally identical sequence are included in the invention. Variants of the nucleic acid polynucleotide sequences, wherein the variants hybridize to at least one disclosed sequence, are considered to be included in the invention. Unique subsequences of the sequences disclosed herein, as determined by, e.g., standard sequence comparison techniques, are also included in the invention.

#### Conservative variations

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Owing to the degeneracy of the genetic code, "silent substitutions" (i.e., substitutions in a nucleic acid sequence which do not result in an alteration in an encoded polypeptide) are an implied feature of every nucleic acid sequence which encodes an amino acid. Similarly, "conservative amino acid substitutions," in one or a few amino acids in an amino acid sequence are substituted with different amino acids with highly similar properties, are also readily identified as being highly similar to a disclosed construct. Such conservative variations of each disclosed sequence are a feature of the present invention.

those nucleic acids which encode identical or essentially identical amino acid sequences, or, where the nucleic acid does not encode an amino acid sequence, to essentially identical sequences. One of skill will recognize that individual substitutions, deletions or additions which alter, add or delete a single amino acid or a small percentage of amino acids (typically less than 5%, more typically less than 4%, 2% or 1%) in an encoded sequence are "conservatively modified variations" where the alterations result in the deletion of an amino acid, addition of an amino acid, or substitution of an amino acid with a chemically similar amino acid. Thus, "conservative variations" of a listed polypeptide sequence of the present invention include substitutions of a small percentage, typically less than 5%, more typically less than 2% or 1%, of the amino acids of the polypeptide sequence, with a conservatively selected amino acid of the same conservative substitution group. Finally, the addition of sequences that do not alter the encoded activity of a nucleic acid molecule, such as the addition of a non-functional sequence, is a conservative variation of the basic nucleic acid.

[0255] Conservative substitution tables providing functionally similar amino acids are well known in the art. The following sets forth example groups which contain natural amino acids that include "conservative substitutions" for one another.

# Conservative Substitution Groups

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1	Alanine (A)	Serine (S)	Threonine (T)	
2	Aspartic acid (D)	Glutamic acid (E)		
3	Asparagine (N)	Glutamine (Q)		
4	Arginine (R)	Lysine (K)		
5	Isoleucine (I)	Leucine (L)	Methionine (M)	Valine (V)
6	Phenylalanine (F)	Tyrosine (Y)	Tryptophan (W)	

# Nucleic Acid Hybridization

[0256] Comparative hybridization can be used to identify nucleic acids of the invention, including conservative variations of nucleic acids of the invention, and this comparative hybridization method is a preferred method of distinguishing nucleic acids of the invention. In addition, target nucleic acids which hybridize to the nucleic acids represented by SEQ ID NO: 3-35 (e.g., 3-19, 20-35, or any other subset of sequences 3-35), 64-85 under high, ultra-high and ultra-ultra high stringency conditions are a feature of the invention. Examples of such nucleic acids include those with one or a few silent or conservative nucleic acid substitutions as compared to a given nucleic acid sequence.

[0257] A test nucleic acid is said to specifically hybridize to a probe nucleic acid when it hybridizes at least ½ as well to the probe as to the perfectly matched complementary target, i.e., with a signal to noise ratio at lest ½ as high as hybridization of the probe to the target under conditions in which the perfectly matched probe binds to the perfectly matched complementary target with a signal to noise ratio that is at least about 5x-10x as high as that observed for hybridization to any of the unmatched target nucleic acids.

Nucleic acids "hybridize" when they associate, typically in solution. Nucleic acids hybridize due to a variety of well characterized physico-chemical forces, such as hydrogen bonding, solvent exclusion, base stacking and the like. An extensive guide to the hybridization of nucleic acids is found in Tijssen (1993) Laboratory Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Acid Probes part I chapter 2, "Overview of principles of hybridization and the strategy of nucleic acid probe

assays," (Elsevier, New York), as well as in Ausubel, *supra*. Hames and Higgins (1995) Gene Probes 1 IRL Press at Oxford University Press, Oxford, England, (Hames and Higgins 1) and Hames and Higgins (1995) Gene Probes 2 IRL Press at Oxford University Press, Oxford, England (Hames and Higgins 2) provide details on the synthesis, labeling, detection and quantification of DNA and RNA, including oligonucleotides.

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[0259] An example of stringent hybridization conditions for hybridization of complementary nucleic acids which have more than 100 complementary residues on a filter in a Southern or northern blot is 50% formalin with 1 mg of heparin at 42°C, with the hybridization being carried out overnight. An example of stringent wash conditions is a 0.2x SSC wash at 65°C for 15 minutes (see, Sambrook, supra for a description of SSC buffer). Often the high stringency wash is preceded by a low stringency wash to remove background probe signal. An example low stringency wash is 2x SSC at 40°C for 15 minutes. In general, a signal to noise ratio of 5x (or higher) than that observed for an unrelated probe in the particular hybridization assay indicates detection of a specific hybridization.

hybridization experiments such as Southern and northern hybridizations are sequence dependent, and are different under different environmental parameters. An extensive guide to the hybridization of nucleic acids is found in Tijssen (1993), *supra*. and in Hames and Higgins, 1 and 2. Stringent hybridization and wash conditions can easily be determined empirically for any test nucleic acid. For example, in determining highly stringent hybridization and wash conditions, the hybridization and wash conditions are gradually increased (e.g., by increasing temperature, decreasing salt concentration, increasing detergent concentration and/or increasing the concentration of organic solvents such as formalin in the hybridization or wash), until a selected set of criteria are met. For example, the hybridization and wash conditions are gradually increased until a probe binds to a perfectly matched complementary target with a signal to noise ratio that is at least 5x as high as that observed for hybridization of the probe to an unmatched target.

[0261] "Very stringent" conditions are selected to be equal to the thermal melting point  $(T_m)$  for a particular probe. The  $T_m$  is the temperature (under defined ionic strength and pH) at which 50% of the test sequence hybridizes to a perfectly matched probe. For the purposes of the present invention, generally, "highly stringent" hybridization and wash

conditions are selected to be about  $5^{\circ}$  C lower than the  $T_m$  for the specific sequence at a defined ionic strength and pH.

"Ultra high-stringency" hybridization and wash conditions are those in which the stringency of hybridization and wash conditions are increased until the signal to noise ratio for binding of the probe to the perfectly matched complementary target nucleic acid is at least 10x as high as that observed for hybridization to any of the unmatched target nucleic acids. A target nucleic acid which hybridizes to a probe under such conditions, with a signal to noise ratio of at least ½ that of the perfectly matched complementary target nucleic acid is said to bind to the probe under ultra-high stringency conditions.

[0263] Similarly, even higher levels of stringency can be determined by gradually increasing the hybridization and/or wash conditions of the relevant hybridization assay. For example, those in which the stringency of hybridization and wash conditions are increased until the signal to noise ratio for binding of the probe to the perfectly matched complementary target nucleic acid is at least 10x, 20X, 50X, 100X, or 500X or more as high as that observed for hybridization to any of the unmatched target nucleic acids. A target nucleic acid which hybridizes to a probe under such conditions, with a signal to noise ratio of at least ½ that of the perfectly matched complementary target nucleic acid is said to bind to the probe under ultra-ultra-high stringency conditions.

[0264] Nucleic acids that do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical. This occurs, *e.g.*, when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code.

#### Unique subsequences

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[0265] In one aspect, the invention provides a nucleic acid that comprises a unique subsequence in a nucleic acid selected from the sequences of O-tRNAs and O-RSs disclosed herein. The unique subsequence is unique as compared to a nucleic acid corresponding to any known O-tRNA or O-RS nucleic acid sequence. Alignment can be performed using, e.g., BLAST set to default parameters. Any unique subsequence is useful, e.g., as a probe to identify the nucleic acids of the invention.

[0266] Similarly, the invention includes a polypeptide which comprises a unique subsequence in a polypeptide selected from the sequences of O-RSs disclosed herein. Here,

the unique subsequence is unique as compared to a polypeptide corresponding to any known polypeptide sequence.

The invention also provides for target nucleic acids which hybridizes under stringent conditions to a unique coding oligonucleotide which encodes a unique subsequence in a polypeptide selected from the sequences of O-RSs wherein the unique subsequence is unique as compared to a polypeptide corresponding to any of the control polypeptides (e.g., parental sequences from which synthetases of the invention were derived, e.g., by mutation). Unique sequences are determined as noted above.

### Sequence comparison, identity, and homology

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[0268] The terms "identical" or percent "identity," in the context of two or more nucleic acid or polypeptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or nucleotides that are the same, when compared and aligned for maximum correspondence, as measured using one of the sequence comparison algorithms described below (or other algorithms available to persons of skill) or by visual inspection.

[0269] The phrase "substantially identical," in the context of two nucleic acids or polypeptides (e.g., DNAs encoding an O-tRNA or O-RS, or the amino acid sequence of an O-RS) refers to two or more sequences or subsequences that have at least about 60%, preferably 80%, most preferably 90-95% nucleotide or amino acid residue identity, when compared and aligned for maximum correspondence, as measured using a sequence comparison algorithm or by visual inspection. Such "substantially identical" sequences are typically considered to be "homologous," without reference to actual ancestry. Preferably, the "substantial identity" exists over a region of the sequences that is at least about 50 residues in length, more preferably over a region of at least about 100 residues, and most preferably, the sequences are substantially identical over at least about 150 residues, or over the full length of the two sequences to be compared.

[0270] For sequence comparison and homology determination, typically one sequence acts as a reference sequence to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are input into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. The sequence comparison algorithm then calculates the percent

sequence identity for the test sequence(s) relative to the reference sequence, based on the designated program parameters.

[0271] Optimal alignment of sequences for comparison can be conducted, e.g., by the local homology algorithm of Smith & Waterman, Adv. Appl. Math. 2:482 (1981), by the homology alignment algorithm of Needleman & Wunsch, J. Mol. Biol. 48:443 (1970), by the search for similarity method of Pearson & Lipman, Proc. Nat'l. Acad. Sci. USA 85:2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by visual inspection (see generally, Ausubel et al., infra).

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[0272]One example of an algorithm that is suitable for determining percent sequence identity and sequence similarity is the BLAST algorithm, which is described in Altschul et al., J. Mol. Biol. 215:403-410 (1990). Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (www.ncbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul et al., supra). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are then extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Cumulative scores are calculated using, for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always > 0) and N (penalty score for mismatching residues; always < 0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, a cutoff of 100, M=5, N=-4, and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength (W) of 3, an expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff & Henikoff (1989) Proc. Natl. Acad. Sci. USA 89:10915).

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In addition to calculating percent sequence identity, the BLAST algorithm also performs a statistical analysis of the similarity between two sequences (*see*, *e.g.*, Karlin & Altschul, Proc. Nat'l. Acad. Sci. USA 90:5873-5787 (1993)). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.1, more preferably less than about 0.01, and most preferably less than about 0.001.

Mutagenesis and Other Molecular Biology Techniques

[0274] General texts which describe molecular biological techniques include Berger and Kimmel, Guide to Molecular Cloning Techniques, Methods in Enzymology volume

15 152 Academic Press, Inc., San Diego, CA (Berger); Sambrook et al., Molecular Cloning - A Laboratory Manual (2nd Ed.), Vol. 1-3, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989 ("Sambrook") and Current Protocols in Molecular Biology, F.M. Ausubel et al., eds., Current Protocols, a joint venture between Greene Publishing Associates, Inc. and John Wiley & Sons, Inc., (supplemented through 1999) ("Ausubel")).

20 These texts describe mutagenesis, the use of vectors, promoters and many other relevant topics related to, e.g., the generation of genes that include selector codons for production of proteins that include unnatural amino acids, orthogonal tRNAs, orthogonal synthetases, and pairs thereof.

[0275] Various types of mutagenesis are used in the invention, e.g., to produce libraries of tRNAs, to produce libraries of synthetases, to insert selector codons that encode unnatural amino acids in a protein or polypeptide of interest. They include but are not limited to site-directed, random point mutagenesis, homologous recombination, DNA shuffling or other recursive mutagenesis methods, chimeric construction, mutagenesis using uracil containing templates, oligonucleotide-directed mutagenesis, phosphorothioate-modified DNA mutagenesis, mutagenesis using gapped duplex DNA or the like, or any combination thereof. Additional suitable methods include point mismatch repair, mutagenesis using repair-deficient host strains, restriction-selection and restriction-

purification, deletion mutagenesis, mutagenesis by total gene synthesis, double-strand break repair, and the like. Mutagenesis, e.g., involving chimeric constructs, are also included in the present invention. In one embodiment, mutagenesis can be guided by known information of the naturally occurring molecule or altered or mutated naturally occurring molecule, e.g., sequence, sequence comparisons, physical properties, crystal structure or the like.

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[0276] The above texts and examples found herein describe these procedures. Additional information is found in the following publications and references cited within: Ling et al., Approaches to DNA mutagenesis: an overview, Anal Biochem. 254(2): 157-178 10 (1997); Dale et al., Oligonucleotide-directed random mutagenesis using the phosphorothioate method, Methods Mol. Biol. 57:369-374 (1996); Smith, In vitro mutagenesis, Ann. Rev. Genet. 19:423-462(1985); Botstein & Shortle, Strategies and applications of in vitro mutagenesis, Science 229:1193-1201(1985); Carter, Site-directed mutagenesis, Biochem. J. 237:1-7 (1986); Kunkel, The efficiency of oligonucleotide 15 directed mutagenesis, in Nucleic Acids & Molecular Biology (Eckstein, F. and Lilley, D.M.J. eds., Springer Verlag, Berlin)) (1987); Kunkel, Rapid and efficient site-specific mutagenesis without phenotypic selection, Proc. Natl. Acad. Sci. USA 82:488-492 (1985); Kunkel et al., Rapid and efficient site-specific mutagenesis without phenotypic selection, Methods in Enzymol. 154, 367-382 (1987); Bass et al., Mutant Trp repressors with new 20 DNA-binding specificities, Science 242:240-245 (1988); Methods in Enzymol. 100: 468-500 (1983); Methods in Enzymol. 154: 329-350 (1987); Zoller & Smith, Oligonucleotidedirected mutagenesis using M13-derived vectors: an efficient and general procedure for the production of point mutations in any DNA fragment, Nucleic Acids Res. 10:6487-6500 (1982); Zoller & Smith, Oligonucleotide-directed mutagenesis of DNA fragments cloned into M13 vectors, Methods in Enzymol. 100:468-500 (1983); Zoller & Smith, 25 Oligonucleotide-directed mutagenesis: a simple method using two oligonucleotide primers and a single-stranded DNA template, Methods in Enzymol. 154:329-350 (1987); Taylor et al., The use of phosphorothioate-modified DNA in restriction enzyme reactions to prepare nicked DNA, Nucl. Acids Res. 13: 8749-8764 (1985); Taylor et al., The rapid generation of oligonucleotide-directed mutations at high frequency using phosphorothioate-modified 30 DNA, Nucl. Acids Res. 13: 8765-8787 (1985); Nakamaye & Eckstein, Inhibition of restriction endonuclease Nci I cleavage by phosphorothioate groups and its application to oligonucleotide-directed mutagenesis, Nucl. Acids Res. 14: 9679-9698 (1986); Sayers et al.,

Y-T Exonucleases in phosphorothioate-based oligonucleotide-directed mutagenesis, Nucl. Acids Res. 16:791-802 (1988); Sayers et al., Strand specific cleavage of phosphorothioatecontaining DNA by reaction with restriction endonucleases in the presence of ethidium bromide, (1988) Nucl. Acids Res. 16: 803-814; Kramer et al., The gapped duplex DNA approach to oligonucleotide-directed mutation construction, Nucl. Acids Res. 12: 9441-5 9456 (1984); Kramer & Fritz Oligonucleotide-directed construction of mutations via gapped duplex DNA, Methods in Enzymol. 154:350-367 (1987); Kramer et al., Improved enzymatic in vitro reactions in the gapped duplex DNA approach to oligonucleotidedirected construction of mutations, Nucl. Acids Res. 16: 7207 (1988); Fritz et al., 10 Oligonucleotide-directed construction of mutations: a gapped duplex DNA procedure without enzymatic reactions in vitro, Nucl. Acids Res. 16: 6987-6999 (1988); Kramer et al., Point Mismatch Repair, Cell 38:879-887 (1984); Carter et al., Improved oligonucleotide site-directed mutagenesis using M13 vectors, Nucl. Acids Res. 13: 4431-4443 (1985); Carter, Improved oligonucleotide-directed mutagenesis using M13 vectors, Methods in Enzymol. 154: 382-403 (1987); Eghtedarzadeh & Henikoff, Use of oligonucleotides to 15 generate large deletions, Nucl. Acids Res. 14: 5115 (1986); Wells et al., Importance of hydrogen-bond formation in stabilizing the transition state of subtilisin, Phil. Trans. R. Soc. Lond. A 317: 415-423 (1986); Nambiar et al., Total synthesis and cloning of a gene coding for the ribonuclease S protein, Science 223: 1299-1301 (1984); Sakamar and Khorana, Total synthesis and expression of a gene for the a-subunit of bovine rod outer segment 20 guanine nucleotide-binding protein (transducin), Nucl. Acids Res. 14: 6361-6372 (1988); Wells et al., Cassette mutagenesis: an efficient method for generation of multiple mutations at defined sites, Gene 34:315-323 (1985); Grundström et al., Oligonucleotide-directed mutagenesis by microscale 'shot-gun' gene synthesis, Nucl. Acids Res. 13: 3305-3316 (1985); Mandecki, Oligonucleotide-directed double-strand break repair in plasmids of 25 Escherichia coli: a method for site-specific mutagenesis, Proc. Natl. Acad. Sci. USA, 83:7177-7181 (1986); Arnold, Protein engineering for unusual environments, Current Opinion in Biotechnology 4:450-455 (1993); Sieber, et al., Nature Biotechnology, 19:456-460 (2001). W. P. C. Stemmer, Nature 370, 389-91 (1994); and, I. A. Lorimer, I. Pastan, Nucleic Acids Res. 23, 3067-8 (1995). Additional details on many of the above methods 30 can be found in Methods in Enzymology Volume 154, which also describes useful controls for trouble-shooting problems with various mutagenesis methods.

[0277] The invention also relates to eukaryotic host cells and organisms for the in vivo incorporation of an unnatural amino acid via orthogonal tRNA/RS pairs. Host cells are genetically engineered (e.g., transformed, transduced or transfected) with the polynucleotides of the invention or constructs which include a polynucleotide of the invention, e.g., a vector of the invention, which can be, for example, a cloning vector or an expression vector. The vector can be, for example, in the form of a plasmid, a bacterium, a virus, a naked polynucleotide, or a conjugated polynucleotide. The vectors are introduced into cells and/or microorganisms by standard methods including electroporation (From et al., Proc. Natl. Acad. Sci. USA 82, 5824 (1985), infection by viral vectors, high velocity ballistic penetration by small particles with the nucleic acid either within the matrix of small beads or particles, or on the surface (Klein et al., Nature 327, 70-73 (1987)).

[0278] The engineered host cells can be cultured in conventional nutrient media modified as appropriate for such activities as, for example, screening steps, activating promoters or selecting transformants. These cells can optionally be cultured into transgenic organisms. Other useful references, e.g. for cell isolation and culture (e.g., for subsequent nucleic acid isolation) include Freshney (1994) Culture of Animal Cells, a Manual of Basic Technique, third edition, Wiley- Liss, New York and the references cited therein; Payne et al. (1992) Plant Cell and Tissue Culture in Liquid Systems John Wiley & Sons, Inc. New York, NY; Gamborg and Phillips (eds) (1995) Plant Cell, Tissue and Organ Culture; Fundamental Methods Springer Lab Manual, Springer-Verlag (Berlin Heidelberg New York) and Atlas and Parks (eds) The Handbook of Microbiological Media (1993) CRC Press, Boca Raton, FL.

Several well-known methods of introducing target nucleic acids into cells are available, any of which can be used in the invention. These include: fusion of the recipient cells with bacterial protoplasts containing the DNA, electroporation, projectile bombardment, and infection with viral vectors (discussed further, below), etc. Bacterial cells can be used to amplify the number of plasmids containing DNA constructs of this invention. The bacteria are grown to log phase and the plasmids within the bacteria can be isolated by a variety of methods known in the art (*see*, for instance, Sambrook). In addition, a plethora of kits are commercially available for the purification of plasmids from bacteria, (see, e.g., EasyPrep<sup>TM</sup>, FlexiPrep<sup>TM</sup>, both from Pharmacia Biotech; StrataClean<sup>TM</sup>, from Stratagene; and, QIAprep<sup>TM</sup> from Qiagen). The isolated and purified plasmids are then further manipulated to produce other plasmids, used to transfect cells or incorporated into

related vectors to infect organisms. Typical vectors contain transcription and translation terminators, transcription and translation initiation sequences, and promoters useful for regulation of the expression of the particular target nucleic acid. The vectors optionally comprise generic expression cassettes containing at least one independent terminator sequence, sequences permitting replication of the cassette in eukaryotes, or prokaryotes, or 5 both, (e.g., shuttle vectors) and selection markers for both prokaryotic and eukaryotic systems. Vectors are suitable for replication and integration in prokaryotes, eukaryotes, or preferably both. See, Giliman & Smith, Gene 8:81 (1979); Roberts, et al., Nature, 328:731 (1987); Schneider, B., et al., Protein Expr. Purif. 6435:10 (1995); Ausubel, Sambrook, Berger (all supra). A catalogue of Bacteria and Bacteriophages useful for cloning is 10 provided, e.g., by the ATCC, e.g., The ATCC Catalogue of Bacteria and Bacteriophage (1992) Gherna et al. (eds) published by the ATCC. Additional basic procedures for sequencing, cloning and other aspects of molecular biology and underlying theoretical considerations are also found in Watson et al. (1992) Recombinant DNA Second Edition Scientific American Books, NY. In addition, essentially any nucleic acid (and virtually any 15 labeled nucleic acid, whether standard or non-standard) can be custom or standard ordered from any of a variety of commercial sources, such as the Midland Certified Reagent Company (Midland, TX mcrc.com), The Great American Gene Company (Ramona, CA available on the World Wide Web at genco.com), ExpressGen Inc. (Chicago, IL available on the World Wide Web at expressgen.com), Operon Technologies Inc. (Alameda, CA) and 20 many others.

### <u>KITS</u>

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[0280] Kits are also a feature of the invention. For example, a kit for producing a protein that comprises at least one unnatural amino acid in a cell is provided, where the kit includes a container containing a polynucleotide sequence encoding an O-tRNA, and/or an O-tRNA, and/or a polynucleotide sequence encoding an O-RS, and/or an O-RS. In one embodiment, the kit further includes at least one unnatural amino acid. In another embodiment, the kit further comprises instructional materials for producing the protein.

#### **EXAMPLES**

[0281] The following examples are offered to illustrate, but not to limit the claimed invention. One of skill will recognize a variety of non-critical parameters that may be altered without departing from the scope of the claimed invention.

# EXAMPLE 1: METHODS OF PRODUCING AND COMPOSITIONS OF AMINOACYLtrna synthetases that incorporate unnatural amino acids in EUKARYOTIC CELLS

[0282] The expansion of the eukaryotic genetic code to include unnatural amino acids with novel physical, chemical or biological properties would provide powerful tools 5 for analyzing and controlling protein function in these cells. Towards this goal, a general approach for the isolation of aminoacyl-tRNA synthetases that incorporate unnatural amino acids with high fidelity into proteins in response to an amber codon in Saccharomyces cerevisiae (S. cerevisiae) is described. The method is based on the activation of GALA responsive reporter genes, HIS3, URA3 or LacZ, by suppression of amber codons between 10 the DNA binding domain and transcriptional activation domain of GAL4. The optimization of a GAL4 reporter for positive selection of active Escherichia coli tyrosyl-tRNA synthetase (EcTyrRS) variants is described. A negative selection of inactive EcTyrRS variants has also been developed with the URA3 reporter by use of a small molecule (5fluroorotic acid (5-FOA)) added to the growth media as a 'toxic allele.' Importantly both 15 positive and negative selections can be performed on a single cell and with a range of stringencies. This can facilitate the isolation of a range of aminoacyl-tRNA synthetase (aaRS) activities from large libraries of mutant synthetases. The power of the method for isolating desired aaRS phenotypes is demonstrated by model selections.

20 The recent addition of unnatural amino acids to the genetic code of [0283] Escherichia coli (E. coli) provides a powerful new approach for analyzing and manipulating protein structure and function both in vitro and in vivo. Amino acids with photoaffinity labels, heavy atoms, keto and olefinic groups and chromophores have been incorporated into proteins in E. coli with an efficiency and fidelity rivaling that of the common twenty amino acids. See, e.g., Chin, et al., (2002), Addition of a Photocrosslinker to the Genetic 25 Code of Escherichia coli, Proc. Natl. Acad.Sci. U. S. A. 99:11020-11024; Chin and Schultz, (2002), In vivo Photocrosslinking with Unnatural Amino Acid Mutagenesis, Chem BioChem 11:1135-1137; Chin et al., (2002), Addition of p-Azido-L-phenylalanine to the Genetic code of Escherichia coli, J. Am. Chem. Soc. 124:9026-9027; Zhang et al., (2002), 30 The selective incorporation of alkenes into proteins in Escherichia coli, Angewandte Chemie. International Ed. in English 41:2840-2842; and, Wang and Schultz, (2002), Expanding the Genetic Code, Chem. Comm. 1-10.

[0284] Unnatural amino acids have been introduced previously into the nicotinic acetylcholine receptor in Xenopus oocytes (e.g., M.W. Nowak, et al. (1998), In vivo incorporation of unnatural amino acids into ion channels in Xenopus oocyte expression system, Method Enzymol. 293:504-529) by microinjection of a chemically misacylated
5 Tetrahymena thermophila tRNA (e.g., M.E. Saks, et al. (1996), An engineered Tetrahymena tRNAGln for in vivo incorporation of unnatural amino acids into proteins by nonsense suppression, J. Biol. Chem. 271:23169-23175, and the relevant mRNA. This has allowed detailed biophysical studies of the receptor in oocytes by the introduction of amino acids containing side chains with unique physical or chemical properties. See, e.g., D.A.
Opin Chem. Biol. 4:645-652. Unfortunately, this methodology is limited to proteins in cells.

Dougherty (2000), Unnatural amino acids as probes of protein structure and function, Curr. Opin. Chem. Biol. 4:645-652. Unfortunately, this methodology is limited to proteins in cells that can be microinjected, and because the tRNA is chemically acylated in vitro, and cannot be re-acylated, the yields of protein are very low. This in turn necessitates sensitive techniques to assay protein function.

[0285] There is interest in the genetic incorporation of unnatural amino acids into 15 proteins in eukaryotic cells in response to an amber codon. See also, H.J. Drabkin et al., (1996), Amber suppression in mammalian cells dependent upon expression of an Escherichia coli aminoacyl-tRNA synthetase gene, Molecular & Cellular Biology 16:907-913; A.K. Kowal, et al., (2001), Twenty-first aminoacyl-tRNA synthetase-suppressor tRNA pairs for possible use in site-specific incorporation of amino acid analogues into proteins in 20 eukaryotes and in eubacteria. [comment], Proc. Natl. Acad. Sci. U. S. A. 98:2268-2273; and, K. Sakamoto, et al., (2002), Site-specific incorporation of an unnatural amino acid into proteins in mammalian cells, Nucleic Acids Res. 30:4692-4699. This would have significant technical and practical advantages, since tRNAs would be re-acylated by their 25 cognate synthetases-leading to large amounts of mutant protein. Moreover, genetically encoded aminoacyl-tRNA synthetases and tRNAs are, in principle, heritable, allowing the unnatural amino acid to be incorporated into proteins through many cell divisions without exponential dilution.

[0286] The steps necessary to add new amino acids to the genetic code of *E. coli* have been described (see, e.g., D.R. Liu, & P.G. Schultz, (1999), Progress toward the evolution of an organism with an expanded genetic code, Proc. Natl. Acad. Sci. U. S. A. 96:4780-4785; and similar principles can be useful for expanding the genetic code of eukaryotes. In the first step, an orthogonal aminoacyl-tRNA synthetase (aaRS)/tRNA<sub>CUA</sub>

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pair is identified. This pair needs to function with the host cells translational machinery, but the aaRS should not charge any endogenous tRNAs with an amino acid and the tRNA<sub>CUA</sub> should not be aminoacylated by any endogenous synthetases. *See, e.g.,* D.R. Liu, et al., *Engineering a tRNA and aminoacyl-tRNA synthetase for the site- specific incorporation of unnatural amino acids into proteins in vivo*, Proc. Natl. Acad. Sci. U. S. A. 94:10092-10097. In a second step, those aaRS/tRNA pairs that are capable of using only the unnatural amino acid are selected from a library of mutant aaRSs. In *E. coli* the selection of unnatural amino acid utilizing variants of MjTyrRS was carried out using two-step 'double sieve' selections. *See, e.g.,* D.R. Liu, & P.G. Schultz, (1999), Progress toward the evolution of an organism with an expanded genetic code, Proc. Natl. Acad. Sci. U. S. A. 96:4780-4785. A modified selection method is used in eukaryotic cells.

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Saccharomyces cerevisiae (S. cerevisiae) was chosen as the eukaryotic host [0287] organism, as it is unicellular, has a rapid generation time, as well as relatively well characterized genetics. See, e.g., D. Burke, et al., (2000) Methods in Yeast Genetics. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY. Moreover, since the translational machinery of eukaryotes is highly conserved (see, e.g., (1996) Translational Control. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY; Y. Kwok, & J.T. Wong, (1980), Evolutionary relationship between Halobacterium cutirubrum and eukaryotes determined by use of aminoacyl-tRNA synthetases as phylogenetic probes, Canadian Journal of Biochemistry 58:213-218; and, (2001) The Ribosome. Cold Spring harbor Laboratory Press, Cold Spring Harbor, NY), it is likely that aaRSs genes for the incorporation of unnatural amino acids discovered in S. cerevisiae can be 'cut and pasted' into higher eukaryotic organisms and used, in partnership with cognate tRNAs (see, e.g., K. Sakamoto, et al., (2002) Site-specific incorporation of an unnatural amino acid into proteins in mammalian cells, Nucleic Acids Res. 30:4692-4699; and, C. Kohrer, et al., (2001), Import of amber and ochre suppressor tRNAs into mammalian cells: a general approach to sitespecific insertion of amino acid analogues into proteins, Proc. Natl. Acad. Sci. U. S. A. 98:14310-14315) to incorporate unnatural amino acids. The expansion of the genetic code of S. cerevisiae is therefore a gateway to expanding the genetic code of complex multicellular eukaryotic organisms. See, e.g., M. Buvoli, et al., (2000), Suppression of nonsense mutations in cell culture and mice by multimerized suppressor tRNA genes, Molecular & Cellular Biology 20:3116-3124. The tyrosyl pair derived from Methanococcus jannaschii TyrRS (MjTyrRS) /tRNA (see e.g., L. Wang, & P.G. Schultz,

(2002), Expanding the Genetic Code, Chem. Comm. 1-10) which was previously used to expand the genetic code of E. coli is not orthogonal in eukaryotic organisms (e.g., P. Fechter, et al., (2001), Major tyrosine identity determinants in Methanococcus jannaschii and Saccharomyces cerevisiae tRNA(Tyr) are conserved but expressed differently, Eur. J.Biochem. 268:761-767) and a new orthogonal pair is required to expand the eukaryotic 5 genetic code. Schimmel and coworkers have shown that the E. coli tyrosyl-tRNA synthetase (EcTyrRS)/tRNA<sub>CUA</sub> pair suppresses amber codons in S. cerevisiae, and that E. coli tRNA<sub>CUA</sub> is not charged by endogenous aminoacyl tRNA synthetases in the yeast cytosol (Figure 2). See also, e.g., H. Edwards, et al., (1991), An Escherichia coli tyrosine transfer RNA is a leucine-specific transfer RNA in the yeast Saccharomyces cerevisiae, 10 Proc. Natl. Acad. Sci. U. S. A. 88:1153-1156; and, H. Edwards, & P. Schimmel (1990), A bacterial amber suppressor in Saccharomyces cerevisiae is selectively recognized by a bacterial aminoacyl-tRNA synthetase, Molecular & Cellular Biology 10:1633-1641. In addition, EcTyrRS has been shown not to charge yeast tRNA in vitro. See, e.g., Y. Kwok, 15 & J.T. Wong, (1980), Evolutionary relationship between Halobacterium cutirubrum and eukaryotes determined by use of aminoacyl-tRNA synthetases as phylogenetic probes, Canadian Journal of Biochemistry 58:213-218; B.P. Doctor, et al., (1966), Studies on the species specificity of yeast and E. coli tyrosine tRNAs, Cold Spring Harbor Symp. Quant. Biol. 31:543-548; and, K. Wakasugi, et al., (1998), Genetic code in evolution: switching species-specific aminoacylation with a peptide transplant, EMBO Journal 17:297-305. 20 Thus, the EcTyrRS /tRNA<sub>CUA</sub> pair is a candidate for an orthogonal pair in S. cerevisiae, as well as in higher eukaryotes (e.g., A.K. Kowal, et al., (2001), Twenty-first aminoacyl-tRNA synthetase-suppressor tRNA pairs for possible use in site-specific incorporation of amino acid analogues into proteins in eukaryotes and in eubacteria.[comment], Proc. Natl. Acad. Sci. U. S. A. 98 (2001) 2268-2273). 25

[0288] To broaden the substrate specificity of EcTyrRS in *E. coli*, Nishimura and coworkers screened an error prone PCR generated library of mutants of EcTyrRS and discovered a mutant with an improved ability to incorporate 3-azatyrosine. *See, e.g.,* F. Hamano-Takaku, et al., (2000), *A mutant Escherichia coli tyrosyltRNA synthetase utilizes the unnatural amino acid azatyrosine more efficiently than tyrosine*, J. Biol. Chem. 275:40324-40328. However, this amino acid is incorporated throughout the proteome of *E. coli*, and the evolved enzyme still prefers tyrosine as a substrate. Yokoyama and coworkers screened a small collection of designed active site variants of EcTyrRS in a wheat germ

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translation system and discovered an EcTyrRS variant that utilizes 3-iodotyrosine more effectively than tyrosine. See, D. Kiga, et al., (2002), An engineered Escherichia coli tyrosyl-tRNA synthetase for site-specific incorporation of an unnatural amino acid into proteins in eukaryotic translation and its application in a wheat germ cellfree system, Proc. Natl. Acad. Sci. U. S. A. 99:9715-9720. In contrast to the enzymes we have evolved in E. 5 coli (e.g., J.W. Chin, et al., (2002), Addition of a Photocrosslinker to the Genetic Code of Escherichia coli, Proc. Natl. Acad. Sci. U. S. A. 99:11020-11024; J.W. Chin, et al., (2002), Addition of p-Azido-L-phenylalanine to the Genetic code of Escherichia coli, J. Am. Chem. Soc. 124:9026-9027; L. Wang, et al., (2001), Expanding the genetic code of Escherichia coli, Science 292:498-500; and, L. Wang, et al., (2002), Adding L-3-(2-naphthyl)alanine to 10 the genetic code of E-coli, J. Am. Chem. Soc. 124:1836-1837), this enzyme still incorporates tyrosine in the absence of the unnatural amino acid. See, e.g., D. Kiga, et al., (2002), An engineered Escherichia coli tyrosyl-tRNA synthetase for site-specific incorporation of an unnatural amino acid into proteins in eukaryotic translation and its application in a wheat germ cellfree system, Proc. Natl. Acad. Sci. U. S. A. 99:9715-9720. 15 Recently, Yokoyama and coworkers have also demonstrated that this EcTyrRS mutant functions with a tRNA<sub>CUA</sub> from Bacillus stearothermophilus to suppress amber codons in mammalian cells. See, K. Sakamoto, et al., (2002), Site-specific incorporation of an unnatural amino acid into proteins in mammalian cells, Nucleic Acids Res. 30:4692-4699.

incorporated with a fidelity similar to that of the common twenty amino acids. To accomplish this goal, a general, in vivo selection method has been used for the discovery of EcTyrRS/tRNA<sub>CUA</sub> variants that function in *S. cerevisiae* to incorporate unnatural amino acids, but none of the common amino acids, in response to the amber codon TAG. A major advantage of a selection is that enzymes which selectively incorporate unnatural amino acids can be rapidly selected and enriched from libraries of 10<sup>8</sup> EcTyrRS active site variants, 6-7 orders of magnitude more diversity than has been screened in vitro. *See, e.g.,* D. Kiga, et al., (2002), *An engineered Escherichia coli tyrosyl-tRNA synthetase for site-specific incorporation of an unnatural amino acid into proteins in eukaryotic translation and its application in a wheat germ cellfree system, Proc. Natl. Acad. Sci. U. S. A. 99:9715-9720. This increase in diversity vastly increases the likelyhood of isolating EcTyrRS variants for the incorporation of a diverse range of useful functionality with very high* 

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fidelity. See, e.g., L. Wang, & P.G. Schultz, (2002), Expanding the Genetic Code, Chem. Comm. 1-10.

[0290] To extend the selection approach to S. cerevisiae, the transcriptional activator protein, GAL4 was used (see Figure 1). See, e.g., A. Laughon, et al., (1984), 5 Identification of two proteins encoded by the Saccharomyces cerevisiae GAL4 gene, Molecular & Cellular Biology 4:268-275; A. Laughon, & R.F. Gesteland, (1984), Primary structure of the Saccharomyces cerevisiae GALA gene, Molecular & Cellular Biology 4:260-267; L. Keegan, et al., (1986), Separation of DNA binding from the transcriptionactivating function of a eukaryotic regulatory protein, Science 231:699-704; and, M. Ptashne, (1988), How eukaryotic transcriptional activators work, Nature 335:683-689. The 10 N-terminal 147 amino acids of this 881 amino acid protein form a DNA binding domain (DBD) that binds DNA sequence specifically. See, e.g., M. Carey, et al., (1989), An aminoterminal fragment of GAL4 binds DNA as a dimer, J. Mol. Biol. 209:423-432; and, E. Giniger, et al., (1985), Specific DNA binding of GALA, a positive regulatory protein of yeast, Cell 40:767-774. The DBD is linked, by an intervening protein sequence, to a C-15 terminal 113 amino acid activation domain (AD) that can activate transcription when bound to DNA. See, e.g., J. Ma, & M. Ptashne, (1987), Deletion analysis of GAL4 defines two transcriptional activating segments, Cell 48:847-853: and, J. Ma, & M. Ptashne, (1987), The carboxy-terminal 30 amino acids of GAL4 are recognized by GAL80, Cell 50:137-142. We envisioned that by placing amber codons towards the N-terminal DBD of a single 20 polypeptide that contained both the N-terminal DBD of GAL4 and its C-terminal AD, amber suppression by the EcTyrRS/tRNA<sub>CUA</sub> pair can be linked to transcriptional activation by GAL4 (Figure 1, Panel A). By the choice of appropriate GAL4 activated reporter genes both positive and negative selections can be performed with the gene (Figure 1, Panel B). While many reporter genes based on complementing the amino acid auxotrophy of a cell 25 can be used for positive selections (eg: URA3, LEU2, HIS3, LYS2), the HIS3 gene is an attractive reporter gene, as the activity of the protein it encodes (imidazole glycerol phosphate dehydratase) can be modulated in a dose dependent manner by addition of 3aminotriazole (3-AT). See, e.g., G.M. Kishore, & D.M. Shah, (1988), Amino acid biosynthesis inhibitors as herbicides, Annual Review of Biochemistry 57:627-663. In S. 30 cerevisiae, fewer genes have been used for negative selections. One of several negative selection strategies (see, e.g., A.J. DeMaggio, et al., (2000), The yeast split-hybrid system, Method Enzymol. 328:128-137; H.M. Shih, et al., (1996), A positive genetic selection for

disrupting protein-protein interactions: identification of CREB mutations that prevent association with the coactivator CBP, Proc. Natl. Acad. Sci. U. S. A. 93:13896-13901; M. Vidal, et al., (1996), Genetic characterization of a mammalian protein-protein interaction domain by using a yeast reverse two-hybrid system. [comment], Proc. Natl. Acad. Sci. U. S. A. 93:10321-10326; and, M. Vidal, et al., (1996), Reverse two-hybrid and one-hybrid 5 systems to detect dissociation of protein-protein and DNA-protein interactions. [comment], Proc. Natl. Acad. Sci. U. S. A. 93:10315-10320) that has been successfully used is the URA3/5-fluroorotic acid (5-FOA) negative selection (e.g., J.D. Boeke, et al., (1984), A positive selection for mutants lacking orotidine-5'-phosphate decarboxylase activity in yeast: 5-fluoroorotic acid resistance, Molecular & General Genetics 197:345-346) system 10 described in the 'reverse two-hybrid' system developed by Vidal and co-workers. See, M. Vidal, et al., (1996), Genetic characterization of a mammalian protein-protein interaction domain by using a yeast reverse two-hybrid system.[comment], Proc. Natl. Acad. Sci. U. S. A. 93:10321-10326; and, M. Vidal, et al., (1996), Reverse two-hybrid and one-hybrid systems to detect dissociation of protein-protein and DNA-protein interactions.[comment], 15 Proc. Natl. Acad. Sci. U. S. A. 93:10315-10320). In the reverse two-hybrid system, a genomically integrated URA3 reporter is placed under a tightly controlled promoter that contains GAL4 DNA binding sites. When two proteins that interact are produced as fusions to the GAL4 DBD and GAL4 AD they reconstitute the activity of GAL4 and activate transcription of URA3. In the presence of 5-FOA, the URA3 gene product converts 20 5-FOA to a toxic product, killing the cell. See, J.D. Boeke, et al., supra. This selection has been used to select for proteins that disrupt a protein-protein interaction and for mutations that disrupt a protein-protein interaction. A variant for screening small molecule inhibitors of protein-protein interactions has also been described. See, e.g., J. Huang, & S.L. Schreiber, (1997) A yeast genetic system for selecting small molecule inhibitors of protein-25 protein interactions in nanodroplets, Proc. Natl. Acad. Sci. U. S. A. 94:13396-13401.

[0291] The appropriate choice of amber codons in full length GAL4 allows efficient positive selections for active EcTyrRS variants using either a HIS3 or URA3 GAL4 activated reporters to complement histidine or uracil auxotrophy in yeast cells. Moreover, the URA3 reporter can be used in negative selections for inactive EcTyrRS variants in the presence of 5-FOA. In addition, colorometric assays using lacZ can be used to read out aminoacyl-tRNA synthetase activity in yeast cells.

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# **RESULTS AND DISCUSSION**

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The EcTyrRS gene was expressed under the control of the constitutive [0292] ADH1 promoter, and the tRNA<sub>CUA</sub> gene was expressed from the same high copy yeast plasmid (pEcTyrRStRNA<sub>CUA</sub>, Figure 1, Panel C). Upon co-transformation of pEcTyrRStRNA<sub>CUA</sub> and a low copy reporter that contains a single amber mutation between 5 the DNA binding domain and activation domain of a chimericGAL4 construct into MaV203, cells grew on selective media lacking histidine and containing 10-20 mM 3-AT (Figure 2). When MaV203 cells were transformed with the same GAL4 construct and either an inactive synthetase mutant (A5) or a construct lacking the EctRNA gene, no growth was observed on 10 mM 3-AT (Figure 2). These experiments establish that 10 EcTyrRS can be constitutively expressed in a functional form from the ADH1 promoter, that there is minimal endogenous amber suppression in MaV203, and that there is little charging of EctRNA<sub>CUA</sub> by yeast synthetases in this system. See, e.g., H. Edwards, et al., (1991), An Escherichia coli tyrosine transfer RNA is a leucine-specific transfer RNA in the yeast Saccharomyces cerevisiae, Proc. Natl. Acad. Sci. U. S. A. 88:1153-1156; and, H. 15 Edwards, & P. Schimmel, (1990), A bacterial amber suppressor in Saccharomyces cerevisiae is selectively recognized by a bacterial aminoacyl-tRNA synthetase, Molecular & Cellular Biology 10:1633-1641. Since EcTyrRS does not charge S. cerevisiae tRNA (e.g., Y. Kwok, & J.T. Wong, (1980), Evolutionary relationship between Halobacterium cutirubrum and eukaryotes determined by use of aminoacyl-tRNA synthetases as 20 phylogenetic probes, Canadian Journal of Biochemistry 58:213-218; B.P. Doctor, et al., (1966), Studies on the species specificity of yeast and E. coli tyrosine tRNAs, Cold Spring Harbor Symp. Quant. Biol. 31:543-548; and, K. Wakasugi, et al., (1998), Genetic code in evolution: switching species-specific aminoacylation with a peptide transplant, EMBO Journal 17:297-305), these experiments confirm that EcTyrRS/EctRNA<sub>CUA</sub> are an 25 orthogonal pair in S. cerevisiae.

[0293] While the first generation GAL4 chimera was able to activate transcription of the weak HIS3 reporter it was unable to activate transcription of the URA3 reporter in MaV203 sufficiently to allow significant growth on concentrations of 3-AT greater than 20 mM, or on – URA plates (Figure 2). For the purposes of selection of EcTyrRS, variants a second generation GAL4 construct was made. This GAL4 reporter was designed to be more active, to have a greater dynamic range, and to avoid the accumulation of revertants. To increase the activity of the GAL4 reporters, full length GAL4 was used (which has a

transcriptional activation activity twice that of a DBD-AD fusion (see, e.g., J. Ma, & M. Ptashne, (1987), Deletion analysis of GALA defines two transcriptional activating segments, Cell 48:847-853) under the control of a strong ADH1 promoter, and a high copy 2-μm plasmid (with a copy number 10-30 times that of the centromeric plasmid of the initial 5 GAL4 chimera) was used. An increase in both the copy number of the plasmid and the activity of the protein it encodes should extend the dynamic range of the reporters. Amber mutations were targeted to the region of the GAL4 gene that encodes amino acid residues 2 and 147 (Figure 3). This region is sufficient for sequence specific DNA binding (see, e.g., M. Carey, et al., (1989), An amino-terminal fragment of GAL4 binds DNA as a dimer, J. Mol. Biol. 209:423-432), and lies to the 5' side of the first cryptic activation domain in the 10 GAL4gene (see, e.g., J. Ma, & M. Ptashne, (1987) Deletion analysis of GAL4 defines two transcriptional activating segments, Cell 48:847-853), such that the truncated products produced in the absence of amber suppression are not anticipated to activate transcription. The choice of amino acid codons to mutate was guided by previous saturation mutagenesis selections on GAL4 (see, e.g., M. Johnston, & J. Dover, (1988), Mutational analysis of the 15 GALA-encoded transcriptional activator protein of Saccharomyces cerevisiae, Genetics 120:63-74), as well as the X-ray structures of the N-terminal DNA binding domain of GALA (see, e.g., R. Marmorstein, et al., (1992), DNA recognition by GALA: structure of a protein-DNA complex. [comment], Nature 356:408-414; and, J.D. Baleja, et al., (1992), Solution structure of the DNA-binding domain of Cd2-GAL4 from S. cerevisiae.[comment], 20 Nature 356:450-453) and the NMR structure of its dimerization region. See, e.g., P. Hidalgo, et al., (2001), Recruitment of the transcriptional machinery through GAL11P: structure and interactions of the GAL4 dimerization domain, Genes & Development 15:1007-1020.

Full length GAL4 was cloned into a small pUC based vector to allow the rapid construction of 10 single amber mutants (at the codons for amino acids L3, I13, T44, F68, R110, V114, T121, I127, S131, T145) by site directed mutageneisis. GAL 4 and the resulting amber mutants were then subcloned into a 2-μm yeast vector under the control of the full length ADH1 promoter to create pGADGAL4 and a series of amber mutants denoted pGADGAL4 (xxTAG) (Figure 1, Panel C), where xx denotes the amino acid codon in the GAL4 gene that was mutated to the amber codon. Each GAL4 mutant was cotransformed with either EcTyrRS/tRNA<sub>CUA</sub> or A5/tRNA<sub>CUA</sub> into MaV203 cells, converting transformants to leucine and tryptophan protrophy. pGADGAL4 itself transformed with

very low efficiency (<10-3 times that of the GAL4 amber mutants) and is presumably deleterious to MaV203 cells at such high copy; no such effect was observed with the amber mutants of GAL4.

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[0295] The phenotypes of GAL4 reporters, in the presence of an active or dead synthetase, were assayed on – URA plates, and 0.1 % 5-FOA plates (Figure 3, Panel A). Five GAL4 mutants (L3TAG, I13TAG, T44TAG, F68TAG, S131TAG) grew on – URA plates and failed to grow on 0.1 % 5-FOA in the presence of either a wild type or inactive EcTyrRS. In these amber mutants, endogenous suppression is apparently sufficient to push the EcTyrRS/tRNA<sub>CUA</sub> mediated suppression beyond the dynamic range of the URA3 reporter in MaV203. Five GAL4 single amber mutants (R110TAG, V114TAG, T121TAG, I127TAG, T145TAG) grew in the absence of uracil and in the presence EcTyrRS/tRNA<sub>CUA</sub>, (but not A5/tRNA<sub>CUA</sub>) and showed the reverse phenotype on 5-FOA. These mutants show EcTyrRS dependent phenotypes that fall within the dynamic range of the URA3 reporter in MaV203. The cleanest EcTyrRS dependent phenotype on both - URA and 0.1% 5-FOA was observed with the R110 TAG mutant of GAL4. However, this mutant showed some blue color in X-GAL assays when cotransformed with A5. To further improve the dynamic range, a series of six double amber mutants of GAL4 were made containing R110 TAG (Figure 3, Panel B), (L3TAG, R110TAG; I13TAG, R110TAG; T44TAG, R110TAG; R110TAG, T121TAG; R110TAG, I127TAG; R110TAG, T145TAG). Four of these double mutants (I13TAG, R110TAG; R110TAG, T121TAG; R110TAG, I127TAG and T145TAG, R110TAG) were unable to grow in the absence of uracil and grew on 0.1 % 5-FOA. These double mutants have activities outside (below) the dynamic range of the plate assays. Two of the double mutants (L3TAG, R110TAG and T44TAG, R110TAG) grew in the presence of wild type EcTyrRS/tRNA<sub>CUA</sub>, but not with A5/tRNA<sub>CUA</sub> on –URA plates; these mutants also showed the expected reciprocal phenotypes on 5-FOA. pGADGAL4 (T44TAG, R110TAG), the more active of these two GAL4 mutants, was selected for a more detailed characterization (Figure 4). MaV203 containing pGADGAL4(T44TAG, R110TAG)/ pEcTyrRS-tRNA<sub>CUA</sub> were blue on X-GAL but the corresponding strain containing pA5/tRNA<sub>CUA</sub> was not. Similarly MaV203 containing pGADGAL4(T44TAG, R110TAG)/pEcTyrRS/tRNA<sub>CUA</sub> grew robustly on plates with 3-AT concentrations up to 75 mM, and on -URA plates but the corresponding strain containing pA5/tRNA<sub>CUA</sub> failed to grow on 10 mM 3AT or in the absence of uracil. Taken together, the EcTyrRS dependent

phenotypes of pGADGAL4(T44TAG, R110TAG) can span the dynamic range of the URA3, HIS3 and lacZ reporters in MaV203.

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It was of interest to determine the activity of GAL4 mutants in which T44 or [0296] R110 were substituted with amino acids other than tyrosine, since the ability to substitute varied amino acids without altering the activity of GAL4 is likely to be useful for selection of mutant aminoacyl-tRNA synthetases that can incorporate unnatural amino acids into proteins. See, e.g., M. Pasternak, et al., (2000), A new orthogonal suppressor tRNA/aminoacyl-tRNA synthetase pair for evolving an organism with an expanded genetic code, Helvetica Chemica Acta 83:2277. A series of five mutants of residue T44 in GAL4, (T44Y, T44W, T44F, T44D, T44K) were constructed in pGADGAL4 (R110TAG), since pGADGAL4 is itself toxic. A similar series of mutants at position R110 in GAL4, (R110Y, R110W, R110F, R110D, R110K) in pGADGAL4(T44TAG) was constructed. These mutants are biased towards the large hydrophobic amino acid side chains that we are interested in incorporating into proteins, but also contain a positively and negatively charged residue as a stringent test of permissiveness. Each mutant was co-transformed with pEcTyrRS/tRNA<sub>CUA</sub> into MaV203 cells and leu+ trp+ isolates assayed for lacZ production by ortho-nitrophenyl-β-D-galactopyranoside (ONPG) hydrolysis (Figure 5). The variation in activity between cells containing GALA with different amino acids substituted for either T44 or R110 was less than 3 fold in all cases. This minimal variability demonstrates the permissiveness of these sites to amino acid substitution without altering the transcriptional activity of GAL4. As expected from the activity of the single amber mutants assayed on selective plates, mutants of T44 made in the GAL4 (R110TAG) background lead to slower hydrolysis of ONPG than mutants of R110 made in the GAL4(T44TAG) background.

[0297] Model enrichment studies were performed to examine the ability of the system to select an active synthetase from a large excess of inactive synthetases (Table 1, Table 2, Figure 6). This selection models the ability to select active synthetases from a library of variants in the presence of an unnatural amino acid. MaV203 cells containing the GAL4(T44, R110) and EcTyrRS/tRNA<sub>CUA</sub> were mixed with a 10 to 10<sup>6</sup> fold excess of GAL4(T44TAG, R110TAG) and A5/tRNA<sub>CUA</sub> as judged by both OD<sub>660</sub>, and the fraction of colonies that turned blue when plated on nonselective – leu,- trp media and assayed by X-GAL overlay. Those cells able to survive on 50 mM 3-AT or in the absence of uracil were selected. The ratio of cells surviving on 3-AT or –URA that were blue in the X-GAL assay to those that were white, when compared to the same ratio in the absence of selection,

clearly demonstrates that the positive selections can enrich active synthetases from dead synthetases by a factor  $>10^5$  (Table 1). Measurement of accurate enrichments for starting ratios greater than  $1:10^5$  was generally not possible, because no more than  $10^6$  cells can be conveniently plated without significant crosstalk between cells leading to unreliable phenotypes.

TABLE 1. MODEL POSITIVE SELECTIONS FOR FUNCTIONAL EcTyrRS.

StartingRatio,EcYRS:A5	1:10	1:10 <sup>2</sup>	1:10 <sup>3</sup>		1:10⁴		1:10 <sup>5</sup>	
Cell Dilution	10 <sup>3</sup>	10 <sup>3</sup>	10²	10 <sup>3</sup>	10	10 <sup>3</sup>	1	10 <sup>3</sup>
-Leu,Trp (#blue <sup>b</sup> )	1360 (81)	1262 (0)	>10 <sup>3</sup> (1)	1774 (0)	>104 (-)	1092 (0)	>104 (-)	1472 (0)
-Ura (#blue <sup>b</sup> )	152 (152)	9 (9)	8 (8)	0 (-)	5 (5)	0 (-)	16 (14)	0 (-)
-His + 50 mM3AT (#blue <sup>b</sup> )	135 (135)	7 (7)	O (-)	0 (-)	3 (3)	0 (-)	10 (10)	0 (-)
Enrichment factor	>10	>10 <sup>2</sup>	>1	10 <sup>3</sup>	>	10 <sup>4</sup>	>	10 <sup>5</sup>

a) Determined by OD<sub>660</sub>

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TABLE 2. MODEL NEGATIVE SELECTIONS FOR NON-FUNCTIONAL EcTyrRS (A5).

StartingRatio, A5: EcYRS	1:10	1:10 <sup>2</sup>	1:10 <sup>3</sup>	1:10	)4
Cell Dilution	10 <sup>3</sup>	10²	10 <sup>2</sup>	10²	10
-Leu,Trp (#white <sup>b</sup> )	353 (22)	1401 (31)	1336 (2)	1375 (0)	>10 <sup>4</sup>
0.1 % 5-FOA (#white <sup>b</sup> )	16 (16)	41 (41)	4 (4)	0 (-)	2 (2)
Enrichment factor	>10	>45	>600	> 0.67×	< 10⁴

a) Determined by OD<sub>660</sub>

[0298] After a positive selection in the presence of unnatural amino acid, the selected cells will contain synthetases able to use natural amino acids and those able to use an added unnatural amino acid. To isolate those synthetases capable of using only the unnatural amino acid, cells encoding synthetases that use natural amino acids must be deleted from the selected clones. This can be accomplished with a negative selection in which the unnatural amino acid is withheld and those synthetases that function with a natural amino acid are removed. A model negative selection was performed in an analogous manner to the model positive selection. EcTyrRS/tRNA<sub>CUA</sub> was mixed with a 10 to 10<sup>5</sup> fold excess of A5/tRNA<sub>CUA</sub> and selection was performed on 0.1 % 5-FOA. Comparison of the ratio of cells surviving on 0.1 % 5-FOA that were white in the X-GAL assay to those that were blue, to the same ratio under non-selective conditions (see Table 2)

b) On X-GAL

b) On X-GAL

makes it clear that the negative selections can enrich dead synthetases from active synthetases by a factor of at least  $0.6 \times 10^4$ . Measurement of accurate enrichments for starting ratios greater than  $1:10^4$  was generally not possible, because no more than  $10^5$  cells could be conveniently plated without significant crosstalk between cells leading to unreliable phenotypes.

[0299] A general approach was developed that allows both positive selection of aaRSs that recognize unnatural amino acids and negative selection of aaRSs that recognize natural amino acids. By varying the stringencies of the selection, a variety of synthetase activities can be isolated. Application of this method to a model selection using variants of EcTyrRS showed enrichments of greater than  $10^5$  in a single round of either positive selection and greater than  $0.6 \times 10^4$  in a single round of negative selection. These observations suggest that this method can provide rapid access to orthogonal aminoacyltRNA synthetases that function to site-specifically incorporation unnatural amino acids with a diversity of side chains into proteins in *S. cerevisiae*. Moreover, enzymes evolved in *S. cerevisiae* can be used in higher eukaryotes.

#### Materials and Methods

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# Vector construction

 $\begin{tabular}{ll} \textbf{[0300]} & The tRNA_{CUA} gene was amplified by PCR using the primers \\ \end{tabular}$ 

tRNA5':GGGGGGACCGGTGGGGGGACCGGTAAGCTTCCCGATAAGGAGCAGG CCAGTAAAAAGCATTACCCCGTGGTGGTTCCCGA (SEQ ID NO:89), and tRNA3':GGCGGCGCTAGCAAGCTTCCCGATAAGGGAGCAGGCCAGTAAAAAGGG AAGTTCAGGGACTTTTGAAAAAAATGGTGGTGGGGGAAGGAT (SEQ ID NO:90) from pESCSU3URA. This, and all other PCR reactions were preformed using the Expand PCR kit from Roche, according to the manufacturers instructions. After restriction endonuclease digestion with NheI and AgeI this tRNA gene was inserted between the same sites in the 2 μm vector pESCTrp (Stratagene) to yield ptRNA<sub>CUA</sub>. The full length ADH1 promoter was amplified by PCR from pDBLeu (Invitrogen) with the primers PADHf: IGGGGGGACCGGTIGGGGGGACCGGTCGGGATCGAAGAAATGATGGTAAATGA AATAGGAAATCAAGG (SEQ ID NO:91) and pADHR:

GGGGGGAATTCAGTTGATTGTATGCTTGGTATAGCTTGAAATATTGTGCAGAA AAAGAAAC (SEQ ID NO:92), digested with AgeI and EcoRI. EcTyrRS was amplified with the primers

pESCTrp1:TCATAACGAGAATTCCGGGATCGAAGAAATGATGGTAAATGAAATA GGAAATCTCATAACGAGAATTCATGGCAAGCAGTAACTTG (SEQ ID NO:93) and pESCTrp2:TTACTACGTGCGGCCGCATGGCAAGCAGTAACTTGTTACTACGTGCG GCCGCTTATTTCCAGCAAATCAGAC (SEQ ID NO:94). The EcTyrRS PCR product were digested with EcoRI and Not I. ptRNA<sub>CUA</sub> was then digested with Age I and Not I. A triple ligation of these three DNAs yielded pEcTyrRS-tRNA<sub>CUA</sub>. Plasmid pA5-tRNA<sub>CUA</sub> in

by overlap PCR using the oligonucleotides F37Afwd:

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alanine) was created

CCGATCGCGCTCGCTTGCGGCTTCGATC (SEQ ID NO:95), N126Afwd:
ATCGCGGCGAACGCCTATGACTGGTTC (SEQ ID NO:96), 182,183,186A,
GTTGCAGGGTTATGCCGCCGCCTGTGCGAACAAACAGTAC (SEQ ID NO:97) and
their reverse complements, as well as the flanking oligonucleotides, 4783:
GCCGCTTTGCTATCAAGTATAAATAG (SEQ ID NO:98), 3256:

which amino acid residues (37,126,182,183 and 186 in the active site are mutated to

- 15 CAAGCCGACAACCTTGATTGG (SEQ ID NO:99) and pEcTyrRS-tRNA<sub>CUA</sub> as a template. The PCR product was digested with EcoRI and Not I and ligated into the large fragment of pEcTyrRS-tRNA<sub>CUA</sub> released upon digestion with the same enzymes. To construct 1st generation DB-AD reporters, the GAL4 DNA binding domain was PCR amplified from pGADT7 (Clontech) using the forward primer pADfwd:
- 20 GGGGACAAGTTTGTACAAAAAAGCAGGCTACGCCAATTTTAATCAAAGTGG
  GAATATTGC (SEQ ID NO:100) or pADfwd(TAG)
  GGGGACAAGTTTGTACAAAAAAGCAGGCTAGGCCAATTTTAATCAAAGTGG
  GAATATTGC (SEQ ID NO:101) and ADrev:
  GGGGACCACTTTGTACAAGAAAGCTGGGTTACTCTTTTTTTGGGTTTGGTGG
- GGTATC (SEQ ID NO:102). These PCR products were cloned into the vector pDEST3-2 (invitrogen) using the Clonase procedure, according to the manufacturer's instructions, yielding pDB-AD and pDB-(TAG)-AD. To construct PGADGAL4 and variants, the GAL4 gene was amplified from pCL1 (Clontech) by PCR using the primers ADH1428-1429

  AAGCTATACCAAGCATACAATC (SEQ ID NO:103), and GAL4C:
- ACAAGGCCTTGCTAGCTTACTCTTTTTTTGGGTTTGGTGGGGTATCTTC (SEQ ID NO:104). This fragment was cloned into the vector pCR2.1 TOPO (Invitrogen) according to the manufacturer's instructions. A clone containing the GAL4 gene (pCR2.1 TOPOGAL4)

was digested with Hind III and the 2.7kb GAL4 fragment gel purified and ligated to the large fragment of pGADT7 that had been digested with Hind III, treated with calf intestinal phosphotase and gel purified. Variants of the GAL4 gene were created by Quikchange reactions (Stratagene), carried out according to the manufacturers instructions, on pCR2.1 using primers listed in the supplementary information. GAL4 mutants were cloned into pGADT7 in the same manner as the wildtype GAL4 gene. All final constructs were confirmed by DNA sequencing.

### Yeast media and manipulations

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S. cerevisiae strain MaV203, (Invitrogen) is MATa; leu2-3,112; trp1 [0301] 109; his  $3\Delta 200$ ; ade 2-101; cvh $2^R$ ; cvh $1^R$ ; GAL4  $\Delta$ ; gal  $80\Delta$ ; GAL1::lacZ; 10 HIS3UASGAL1::HIS3@LYS2; SPAL10UASGAL1::URA3. Yeast media were purchased from Clontech, 5-FOA and X-GAL were from Invitrogen and 3-AT was from BIO 101. YPER (Yeast Protein Extraction Reagent) and ONPG were purchased from Pierce Chemicals. Plasmid transformations were performed by the PEG/Lithium actetate method 15 (see, e.g., D. Burke, et al., (2000) Methods in Yeast Genetics. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY) and transformants selected on the appropriate synthetic complete dropout media. To test the phenotypes conferred by various plasmid combinations on MaV203 yeast colonies from synthetic complete dropout plates of each transformation were resuspended in 15  $\mu$ L of sterile water and streaked on the selective 20 media of interest. Each phenotype was confirmed with at least five independent colonies. X-GAL assays were performed by the agarose overlay method. See, I.G. Serebriiskii, & E.A. Golemis, (2000), Uses of lacZ to study gene function: evaluation of beta-galactosidase assays employed in the yeast two-hybrid system, Analytical Biochemistry 285:1-15. Briefly, colonies or cell patches were lysed on agar plates by several additions of 25 neatchlorofom. After chloroform evaporation 1 % agarose containing 0.25g/L of XGAL and buffered with 0.1 M Na<sub>2</sub>PO<sub>4</sub> was applied to the plate surface. Once the agarose was set, plates were incubated at 37 °C for 12 h. ONPG assays were carried out by inoculation of 1 mL of SD-leu,-trp in a 96 well block with a single colony and incubated at 30°C with shaking. The  $OD_{660}$  of  $100\mu$ L of cells, and several dilutions of cells were recorded in 30 parallel in a 96 well microtiter plate. Cells (100  $\mu$ L) were mixed with a 100 $\mu$ L of YPER:ONPG (1X PBS, 50% v/v YPER, 20mM MgCl<sub>2</sub>, 0.25 % v/v β-mercaptoethanol, and 3 mM ONPG) and incubated with shaking at 37°C. Upon color development, cells were pelleted by centrifugation, the supernatant transferred to a clean 96 well microtiter plate

(Nunclon, cat.# 167008), and the A420 recorded. All data shown are the mean of trials from at least 4 independent clones and the error bars shown represent the standard deviation. ONPG hydrolysis was calculated using the equation: beta-galactosidase units= 1000. A420/(V.t.OD<sub>660</sub>), where V is the volume of cells in milliliters, t is the time of incubation in minutes. See, e.g., I.G. Serebriiskii, & E.A. Golemis, (2000), Uses of lacZ to study gene function: evaluation of beta-galactosidase assays employed in the yeast two-hybrid system, Analytical Biochemistry 285:1-15. One beta-galactosidase unit corresponds to the hydrolysis of 1  $\mu$ mol of ONPG per minute per cell. See, Serebriiskii and Golemis, supra. Spectrophotometric readings were performed on a SPECTRAmax 190 plate reader.

## **Model Selections**

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Positive selections: Two overnight cultures were grown in SD -Leu, -Trp. [0302] One contained MaV203 harboring pEcTyrRS-tRNA<sub>CUA</sub>/ pGADGAL4(T44, R110TAG) and the other pA5-tRNASU3/pGADGAL4(T44, R110TAG). These cells were harvested by centrifugation and resuspended in 0.9 % NaCl by vortexing. The two cell solutions were then diluted to identical OD660s. MaV203 harboring pEcTyrRS-tRNA<sub>CUA</sub>/ pGADGAL4(T44,R110TAG) were serially diluted over 7 orders of magnitude and each dilution was then mixed 1:1 vol:vol with undiluted MaV203 harboring pA5-tRNA<sub>CUA</sub>/ pGADGAL4(T44, R110TAG) to afford defined ratios of cells containing active and inactive tyrosyl-tRNA synthetase. For each ratio a second serial dilution was perfomed in which the number of cells was decreased but the ratio of cells harboring pEcTyrRStRNA<sub>CUA</sub>/ pGADGAL4(T44,R110TAG) and pA5-tRNA<sub>CUA</sub>/ pGADGAL4(T44,R110TAG) was maintained. These dilutions were plated on SD -Leu, -trp, SD -Leu,-Trp,- URA and SD - Leu, - Trp, -His + 50mM 3-AT. After 60 h the number of colonies on each plate was counted, using an Eagle Eye CCD camera (Stratagene), and the phenotype of survivors were confirmed with a X-GAL beta-galactosidase assay. Cells from several individual blue or white colonies were isolated and grown to saturation in SD -leu,-trp and the plasmid DNA isolated by standard methods. The identity of the EcTyrRS variant was confirmed by DNA sequencing.

[0303] Negative selection: The model negative selection was performed in an analogous manner to the positive selection except that MaV203 harboring pA5-tRNA<sub>CUA</sub>/pGADGAL4(T44,R110TAG) were serially diluted and mixed with a fixed density of MaV203 harboring pEcTyrRS-tRNA<sub>CUA</sub>/ pGADGAL4(T44,R110TAG). Cells

were plated on SD –leu, -trp +0.1 % 5-FOA, the number of colonies counted after 48 hours and the plates processed as described above.

[0304] The following oligonucleotides (Table 3) were used in combination with their reverse complements to construct site-directed mutants by Quikchange mutagenesis.

5 The position of the mutation is denoted by bold text.

# TABLE 3: OLIGONUCLEOTIDES USED TO CONSTRUCT SITE-DIRECTED MUTANTS.

Amber

Mutants Oligo Sequence

- 10 L3TAG 5'-ATGAAGTAGCTGTCTTCTATCGAACAAGCATGCG-3' (SEQ ID NO:66) I13TAG 5'-CGAACAAGCATGCGATTAGTGCCGACTTAAAAAG-3' (SEQ ID NO:67) T44TAG 5'-CGCTACTCTCCCAAATAGAAAAGGTCTCCGCTG-3' (SEQ ID NO:68) F68TAG 5'-CTGGAACAGCTATAGCTACTGATTTTTCCTCG-3' (SEQ ID NO:69) R110TAG 5'-GCCGTCACAGATTAGTTGGCTTCAGTGGAGACTG-3' (SEQ ID
- 15 NO:70)
  V114TAG 5'-GATTGGCTTCATAGGAGACTGATATGCTCTAAC-3' (SEQ ID NO:71)
  T121TAG 5'-GCCTCTATAGTTGAGACAGCATAGAATAATGCG-3' (SEQ ID NO:72)
  I127TAG 5'-GAGACAGCATAGATAGAGTGCGACATCATCATCGG-3' (SEQ ID NO:73)
- 20 S131TAG 5'-GAATAAGTGCGACATAGTCATCGGAAGAGAGTAGTAG-3' (SEQ ID NO:74)
  T145TAG 5'-GGTCAAAGACAGTTGTAGGTATCGATTGACTCGGC-3' (SEQ ID NO:75)
  Permissive
- 25 Site Mutants Oligo Sequence T44F 5'-CGCTACTCTCCCCAAATTTAAAAAGGTCTCCGCTG-3' (SEQ ID NO:76) T44Y 5'-CGCTACTCTCCCCAAATATAAAAAGGTCTCCGCTG-3' (SEQ ID NO:77) T44W 5'-CGCTACTCTCCCCAAATGGAAAAAGGTCTCCGCTG-3' (SEQ ID NO:78) T44D 5'-CGCTACTCTCCCCAAAGATAAAAAGGTCTCCGCTG-3' (SEQ ID NO:79)
- T44K 5'-CGCTACTCTCCCCAAAAAAAAAAAAGGTCTCCGCTG-3' (SEQ ID NO:80)
  R110F 5'-GCCGTCACAGATTTTTTGGCTTCAGTGGAGACTG-3' (SEQ ID NO:81)
  R110Y 5'-GCCGTCACAGATTATTTGGCTTCAGTGGAGACTG-3' (SEQ ID NO:82)
  R110W 5'-GCCGTCACAGATTGGTTGGCTTCAGTGGAGACTG-3' (SEQ ID NO:83)
  R110D 5'-GCCGTCACAGATGATTTGGCTTCAGTGGAGACTG-3' (SEQ ID NO:84)
- 35 R110K 5'-GCCGTCACAGATAAATTGGCTTCAGTGGAGACTG-3' (SEQ ID NO:85)

### **EXAMPLE 2: AN EXPANDED EUKARYOTIC GENETIC CODE**

- [0305] A general and rapid route for the addition of unnatural amino acids to the genetic code of *Saccharomyces cerevisiae* is described. Five amino acids have been incorporated into proteins efficiently, with high fidelity, in response to the nonsense codon
- 40 TAG. The side chains of these amino acids contain a keto group, which can be uniquely modified in vitro and in vivo with a wide range of chemical probes and reagents; a heavy

atom-containing amino acid for structural studies; and photocrosslinkers for cellular studies of protein interactions. This methodology not only removes the constraints imposed by the genetic code on our ability to manipulate protein structure and function in yeast, it provides a gateway to the systematic expansion of the genetic codes of multicellular eukaryotes.

- Although chemists have developed a powerful array of methods and strategies to synthesize and manipulate the structures of small molecules (*see, e.g.*, E. J. Corey, & X.-M. Cheng, The Logic of Chemical Synthesis (Wiley-Interscience, New York, 1995)), the ability to rationally control protein structure and function is still in its infancy. Mutagenesis methods are limited to the common 20 amino acid building blocks, although in a number of cases it has been possible to competitively incorporate close structural analogues of common amino acids throughout the proteome. *See, e.g.*, K. Kirshenbaum, et al., (2002), ChemBioChem 3:235-7; and, V. Doring et al., (2001), Science 292:501-4. Total synthesis (*see, e.g.*, B. Merrifield, (1986), Science 232:341-7 (1986)), and semi-synthetic methodologies (*see, e.g.*, D. Y. Jackson et al., (1994) Science 266:243-7; and, P. E.
- Dawson, & S. B. Kent, (2000), <u>Annual Review of Biochemistry</u> 69:923-60, have made it possible to synthesize peptides and small proteins, but have more limited utility with proteins over 10 kilodaltons (kDa). Biosynthetic methods that involve chemically acylated orthogonal tRNAs (*see*, *e.g.*, D. Mendel, et al., (1995), <u>Annual Review of Biophysics and Biomolecular Structure</u> 24:435-462; and, V. W. Cornish, et al. (Mar. 31, 1995),
- Angewandte Chemie-International Edition in English 34:621-633) have allowed unnatural amino acids to be incorporated into larger proteins, both *in vitro* (*see*, *e.g.*, J. A. Ellman, et al., (1992), Science 255:197-200) and in microinjected cells (see, e.g., *see*, *e.g.*, D. A. Dougherty, (2000), Current Opinion in Chemical Biology 4:645-52). However, the stoichiometric nature of chemical acylation severely limits the amount of protein that can be generated. Thus, despite considerable efforts the properties of proteins, and possibly entire organisms, have been limited throughout evolution by the twenty genetically encoded amino acids (with the rare exceptions of pyrrolysine and selenocysteine (*see*, *e.g.*, A. Bock et al., (1991), Molecular Microbiology 5:515-20; and, G. Srinivasan, et al., (2002), Science 296:1459-62)).
- 30 [0307] To overcome this limitation, new components were added to the protein biosynthetic machinery of the prokaryote *Escherichia coli* (*E. coli*) (e.g., L. Wang, et al., (2001), <u>Science</u> 292:498-500), which make it possible to genetically encode unnatural amino acids *in vivo*. A number of new amino acids with novel chemical, physical or

biological properties have been incorporated efficiently and selectively into proteins in response to the amber codon, TAG. See, e.g., J. W. Chin et al., (2002), Journal of the American Chemical Society 124:9026-9027; J. W. Chin, & P. G. Schultz, (2002), ChemBioChem 11:1135-1137; J. W. Chin, et al., (2002), PNAS United States of America 99:11020-11024: and, L. Wang, & P. G. Schultz, (2002), Chem. Comm., 1:1-10. However, because the translational machinery is not well conserved between prokaryotes and eukaryotes, components of the biosynthetic machinery added to E.coli cannot generally be used to site-specifically incorporate unnatural amino acids into proteins to study or manipulate cellular processes in eukaryotic cells.

10 [0308] Thus, translational components were created that would expand the number of genetically encoded amino acids in eukaryotic cells. Saccharomyces cerevisiae was chosen as the initial eukaryotic host organism, because it is a useful model eukaryote, genetic manipulations are facile (see, e.g., D. Burke, et al., (2000), Methods in Yeast Genetics (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY), and its translational machinery is highly homologous to that of higher eukaryotes (see, e.g., T. R. 15 Hughes, (2002), Funct. Integr. Genomics 2:199-211). The addition of new building blocks to the S. cerevisiae genetic code requires a unique codon, tRNA, and aminoacyl-tRNA synthetase ('aaRS') that do not cross-react with any components of the yeast translational machinery (see, e.g., Noren et al., (1989) Science 244:182; Furter (1998) Protein Sci. 7:419; and, Liu et al., (1999) PNAS USA 96:4780). One candidate orthogonal pair is the amber 20 suppressor tyrosyl-tRNA synthetase-tRNA<sub>CUA</sub> pair from E. coli (see, e.g., H. M. Goodman, et al., (1968), Nature 217:1019-24; and, D. G. Barker, et al., (1982), FEBS Letters 150:419-23), E. coli tyrosyl-tRNA synthetase (TyrRS) efficiently aminoacylates E. coli tRNA<sub>CUA</sub> when both are genetically encoded in S. cerevisiae but does not aminoacylate S. cerevisiae 25 cytoplasmic tRNAs. See, e.g., H. Edwards, & P. Schimmel, (1990), Molecular & Cellular Biology 10:1633-41; and, H. Edwards, et al., (1991), PNAS United States of America 88:1153-6. In addition, E. coli tyrosyl tRNA<sub>CUA</sub> is a poor substrate for S. cerevisiae aminoacyl-tRNA synthetases (see, e.g., V. Trezeguet, et al., (1991), Molecular & Cellular Biology 11:2744-51.) but is processed and exported from the nucleus to the cytoplasm (see, 30 e.g., S. L. Wolin, & A. G. Matera, (1999) Genes & Development 13:1-10) and functions efficiently in protein translation in S. cerevisiae. See, e.g., H. Edwards, & P. Schimmel, (1990) Molecular & Cellular Biology 10:1633-41; H. Edwards, et al., (1991), PNAS United States of America 88:1153-6; and, V. Trezeguet, et al., (1991), Molecular & Cellular

Biology 11:2744-51. Moreover, *E. coli* TyrRS does not have an editing mechanism and therefore should not proofread an unnatural amino acid ligated to the tRNA.

[0309] To alter the amino acid specificity of the orthogonal TyrRS so that it aminoacylates tRNA<sub>CUA</sub> with a desired unnatural amino acid and none of the endogenous amino acids, a large library of TyrRS mutants was generated and subject to a genetic 5 selection. On the basis of the crystal structure of the homologous TyrRS from Bacillus stearothermophilus (see, e.g., P.Brick, et al., (1989), Journal of Molecular Biology 208:83) five residues ((B. stearothermophilus, Figure. 7, Panel A)) in the active site of E coli. TyrRS that are within 6.5 Å of the para position of the aryl ring of bound tyrosine were mutated. For example, to create the EcTyrRS library of mutants the five positions targeted 10 for mutation were first converted to alanine codons to produce the A5RS gene. This was split between two plasmids at a unique Pst I site in the gene. The library was created essentially as described by techniques known in the art (see, e.g., Stemmer et al., (1993) Biotechniques 14:256-265). One plasmid contains the 5' half of the A5RS gene, the other 15 plasmid contains the 3' half of the A5RS gene. Mutagenesis was performed on each fragment by PCR with oligonucleotide primers for the amplification of the whole plasmid. The primers are doped, containing NNK (N = A + G + T + C and K = G + T) and Bsa I restriction endonuclease recognition sites. Digestion with Bsa I and ligation yielded two circular plasmids, each containing mutant copies of one half of the EcTyrRS gene. The two 20 plasmids were then digested with Pst I and assembled into a single plasmid by ligation, leading to assembly of the full-length mutant genes. The mutant EcTyrRS genes were excised from this plasmid and ligated into pA5RS/tRNA<sub>CUA</sub> between EcoR I and Not I sites. The library was transformed into S. cerevisiae Mav203: pGADGAL4 (2TAG) using the PEG-lithium acetate method yielding  $\sim 10^8$  independent transformants.

[0310] A selection strain of S. cerevisiae [MaV203: pGADGALA (2 TAG) (see, e.g., M. Vidal, et al., (1996), PNAS United States of America 93:10321-6; M. Vidal, et al., (1996), PNAS United States of America 93:10315-20l and, Chin et al., (2003) Chem. Biol. 10:511)] was transformed with the library to afford 10<sup>8</sup> independent transformants and grown in the presence of 1 mM unnatural amino acid (Figure 8, Panel C). Suppression of two permissive amber codons in the transcriptional activator GALA leads to the production of full-length GALA and the transcriptional activation of the GALA-responsive HIS3, URA3, and lacZ reporter genes (Figure 8, Panel A). For example, the permissive cdons are for T44 and R110 of Gal4. Expression of HIS3 and URA3 in media lacking uracil (-ura),

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or containing 20 mM 3-aminotriazole (*see*, *e.g.*, G. M. Kishore, & D. M. Shah, (1988), Annual Review of Biochemistry 57, 627-63) (3-AT, a competitive inhibitor of the His3 protein) and lacking histidine (– his), allows clones expressing active aaRS-tRNA<sub>CUA</sub> pairs to be positively selected. If a mutant TyrRS charges the tRNA<sub>CUA</sub> with an amino acid, then the cell biosynthesizes histidine and uracil and survives. Surviving cells were amplified in the absence of 3-AT and unnatural amino acid to remove full-length GAL4 from cells that selectively incorporate the unnatural amino acid. To remove clones that incorporate endogenous amino acids in response to the amber codon, cells were grown on media containing 0.1% 5-fluorootic acid (5-FOA) but lacking the unnatural amino acid. Those cells expressing URA3, as a result of suppression of the GAL4 amber mutations with natural amino acids, convert 5-FOA to a toxic product, killing the cell. *See*, *e.g.*, J. D. Boeke, et al., (1984), Molecular & General Genetics 197:345-6. Surviving clones were amplified in the presence of unnatural amino acid and reapplied to the positive selection. The lacZ reporter allows active and inactive synthetase-tRNA pairs to be discriminated colorometrically (Figure 8, Panel B).

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With the use of this approach, five novel amino acids with distinct steric and [0311] electronic properties (Figure 7, Panel B) were independently added to the genetic code of S. cerevisiae. These amino acids include p-acetyl-L-phenylalanine (1), p-benzoyl-Lphenylalanine (2), p-azido-L-phenylalanine (3), O-methyl-L-tyrosine (4), and p-iodo-Lphenylalanine (5) (indicated by the numbers in Figure 7, Panel B). The unique reactivity of the keto functional group of p-acetyl-L-phenylalanine allows selective modification of proteins with an array of hydrazine- or hydroxylamine-containing reagents in vitro and in vivo (See, e.g., V. W. Cornish, et al., (Aug. 28, 1996), Journal of the American Chemical Society 118:8150-8151; and, Zhang, Smith, Wang, Brock, Schultz, in preparation). The heavy atom of p-iodo-L-phenylalanine can prove useful for phasing x-ray structure data (with the use of multiwavelength anomalous diffraction). The benzophenone and phenylazide side chains of p-benzoyl-L-phenylalanine and p-azido-L-phenylalanine allow efficient in vivo and in vitro photocrosslinking of proteins (see e.g., Chin et al., (2002) J. Am. Chem. Soc., 124:9026; Chin and Schultz, (2002) Chem. Bio. Chem. 11:1135; and, Chin et al., (2002) PNAS,USA 99:11020). The methyl group of O-methyl-L-tyrosine can be readily substituted with an isotopically labeled methyl group as a probe of local structure and dynamics with the use of nuclear magnetic resonance and vibrational spectroscopy. After three rounds of selection (positive-negative-positive), several colonies were isolated

whose survival on –ura or on 20 mM 3-AT –his media was strictly dependent on the addition of the selected unnatural amino acid. See, Figure 8, Panel D. The same clones were blue on x-gal only in the presence of 1 mM unnatural amino acid. These experiments demonstrate that the observed phenotypes result from the combination of the evolved aminoacyl–tRNA synthetase–tRNA<sub>CUA</sub> pairs and their cognate amino acids (see, Table 4).

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For example, to select mutant synthetases, cells (~ 109).were grown for 4 [0312] hours in liquid SD -leu, -trp + 1 mM amino acid. Cells were then harvested by centrifugation, resuspended in 0.9 % NaCl, and plated on SD -leu, -trp, -his + 20 mM 3-AT, + 1mM unnatural amino acid or SD -leu, -trp, -ura, + 1 mM unnatural amino acid. After 48 to 60 hours at 30°C the cells were scraped from the plates into liquid SD -leu, -trp and grown for 15 hours at 30°C. Cells were harvested by centrifugation, resuspended in 0.9 % NaCl and plated on SD -leu, -trp + 0.1 % 5-FOA. After 48 hours at 30°C cells were scraped into liquid SD -leu, -trp + 1 mM unnatural amino acid and grown for 15 hours. Cells were then harvested by centrifugation, resuspended in 0.9 % NaCl, and plated on SD -leu, -trp, his + 20 mM 3-AT, + 1 mM unnatural amino acid or SD -leu, -trp, -ura, + 1 mM unnatural amino acid. To screen phenotypes of selected cells, colonies (192) from each selection were transferred to wells of 96 well blocks containing 0.5 mL of SD -leu, -trp and grown at 30°C for 24 hours. Glycerol (50 % v/v; 0.5 mL) was added to each well, and the cells replica plated onto agar (SD -leu, -trp; SD -leu, -trp, -his, + 20 mM 3-AT; SD -leu, -trp, ura) in the presence or absence of 1 mM unnatural amino acid. X-Gal assays were performed on SD -leu, -trp plates using the agarose overlay method.

[0313] To further demonstrate that the observed phenotypes are due to site-specific incorporation of the unnatural amino acids by the orthogonal mutant TyrRS/tRNA pairs, mutants of human superoxide dismutase 1 (hSOD) (see, e.g., H. E. Parge, et al., (1992), PNAS United States of America 89:6109-13) containing each unnatural amino acid were generated and characterized.

[0314] For example, the addition of DNA encoding a C-terminal hexahistidine tag, and mutation of the codon for Trp 33 to an amber codon in the human superoxide dismutase gene was performed by overlap PCR using PS356 (ATCC) as a template. hSOD (Trp 33 TAG) HIS was cloned between the GAL1 promoter and CYC1 terminator from pYES2.1 (Invitrogen, Carlsbad, CA USA). Mutant synthetase and tRNA genes on pECTyrRS-tRNA<sub>CUA</sub> derived plasmids were co-transformed with pYES2.1 hSOD (Trp 33 TAG) HIS

into the strain InvSc (Invitrogen). For protein expression, cells were grown in SD -trp, -ura + raffinose and expression induced at an OD<sub>660</sub> of 0.5 by the addition of galactose. HSOD mutants were purified by Ni-NTA chromatography (Qiagen, Valencia, CA, USA).

[0315] Production of hexa-histidine-tagged hSOD from a gene containing an amber codon at position 33 was strictly dependent on *p*-acetylPheRS-1–tRNA<sub>CUA</sub> and 1 mM *p*-acetyl-L-phenylalanine (<0.1% by densitometry, in the absence of either component) (See Figure 9). *p*-Acetyl-L-phenylalanine containing full-length hSOD was purified (e.g., by Ni-NTA affinity chromatography) with a yield of 50 ng/mL, comparable to that purified from cells containing *E. coli* TyrRStRNA<sub>CUA</sub>. For comparison, wild type hSODHIS could be purified with a yield of 250 ng/mL under identical conditions.

[0316] Figure 9 illustrates protein expression of hSOD (33TAG)HIS in S. cerevisiae genetically encoding unnatural amino acids (as illustrated in Figure 7, Panel B and indicated in Figure 9 by their numbering in Figure 7, Panel B). The top portion of Figure 9 illustrates SDS-polyacrylamide gel electrophoresis of hSOD purified from yeast in the presence (+) and absence (-) of the unnatural amino acid indicated by the number, which corresponds to unnatural amino illustrated in Figure 7, Panel B stain with Coomassie. Cells contain the mutant synthetase-tRNA pair selected for the amino acid indicated. The center portion of Figure 9 illustrates a western blot probed with an antibody against hSOD. The bottom portion of Figure 9 illustrates a western blot probed with an antibody against the C-terminal His6 tag.

The identity of the amino acid incorporated was determined by subjecting a tryptic digest of the mutant protein to liquid chromatography and tandem mass spectrometry. For example, for mass spectrometry protein bands were visualized by colloidal Coomassie stain. Gel bands corresponding to wild-type and mutant SOD were excised from polyacrylamide gels, sliced into 1.5-mm cubes, reduced and alkylated, then subjected to trypsin hydrolysis essentially as described. *See*, *e.g.*, A. Shevchenko, et al., (1996), Analytical Chemistry 68, 850-858. Tryptic peptides containing the unnatural amino acid were analyzed by nanoflow reversed-phase HPLC/μESI/MS with an LCQ ion trap mass spectrometer. Liquid chromatography tandem mass spectrometry (LC-MS/MS) analysis was performed on a Finnigan LCQ Deca ion trap mass spectrometer (Thermo Finnigan) fitted with a Nanospray HPLC (Agilent 1100 series). *See*, *e.g.*, **Figure 10**, **Panels A-H.** 

The precursor ions corresponding to the singly and doubly charged ions of [0318] the peptide Val-Y\*-Gly-Ser-Ile-Lys (SEQ ID NO:87) containing the unnatural amino acids (denoted Y\*) were separated and fragmented with an ion trap mass spectrometer. The fragment ion masses could be unambiguously assigned, confirming the site-specific incorporation of p-acetyl-L-phenylalanine (see, Figure 10, Panel A). No indication of tyrosine or other amino acids in place of p-acetyl-L-phenylalanine was observed, and a minimum of 99.8% incorporation purity was obtained from the signal-to-noise ratio of the peptide spectra. Similar fidelity and efficiency in protein expression were observed when pbenzoylPheRS-1, p-azidoPheRS-1, O-meTyrRS-1, or p-iodoPheRS-1 was used to incorporate p-benzoyl-L-phenylalanine, p-azido-L-phenylalanine, O-methyl-L-tyrosine, or 10 p-iodo-L-phenylalanine into hSOD (See, Figure 9, and Figure 10, Panels A-H). In the experiments, p-Azido-L-phenylalanine is reduced to p-amino-L-phenylalanine in sample preparation, and the latter is observed in mass spectra. The reduction does not occur in vivo by chemical deriviatation of purified SOD containing p-azido-L-phenylalanine. In control experiments, hexa-histidine-tagged hSOD containing trypotophan, tyrosine, and leucine at 15 position 33 was prepared and subject to mass spectrometry (See, Figure 10, Panels F, G and H). Ions containing amino acid 33 were clearly visible in the mass spectra of these samples.

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The independent addition of five unnatural amino acids to the genetic code [0319] of S. cerevisiae demonstrates the generality of our method and suggests that it can be applicable to other unnatural amino acids including spin-labeled, metal-binding, or photoisomerizable amino acids. This methodology can allow the generation of proteins with new or enhanced properties as well as facilitate control of protein function in yeast. Moreover, in mammalian cells the E. coli tyrosyl-tRNA synthetase forms an orthogonal pair with the B. stearothermophilus tRNA<sub>CUA</sub>. See, e.g., Sakamoto et al., (2002) Nucleic Acids Res. 30:4692. Therefore one can use the aminoacyl-tRNA synthethases that have been evolved in yeast to add unnatural amino acids to the genetic codes of higher eukaryotes.

TABLE 4. SEQUENCES OF SELECTED AMINOACYL-TRNA SYNTHETASES.

Residue #	37	126	182	183	186	# clones
Ec TyrRS	Tyr	Asn	Asp	Phe	Leu	

			<del></del>			
p-IodoPheRS-1	Val	Asn	Ser	Tyr	Leu	1/8
p-IodoPheRS-2	Ile	` Asn	Ser	Met	Leu	1/8
p-IodoPheRS-3	Val	Asn	Ser	Met	Ala	6/8
p-OMePheRS-1	Val	Asn	Ser	Met	Leu	5/13
p-OMePheRS-2	Thr	Asn	Thr	Met	Leu	1/13
p-OMePheRS-3	Thr	Asn	Thr	Tyr	Leu	1/13
p-OMePheRS-4	Leu	Asn	Ser	Met	Ser	1/13
p-OMePheRS-5	Leu	Asn	Ser	Met	Ala	1/13
p-OMePheRS-6	Thr	Asn	Arg	Met	Leu	4/13
p-acetylPheRS-1 <sup>a</sup>	Ile	Asn	Gly	Met	Ala	10/10
p-benzoylPheRS-1	Gly	Asn	Gly	Phe	Ala	1/2
p-benzoylPheRS-2	Gly	Asn	Gly	Tyr	Met	1/2
p-azidoPheRS-1	Leu	Asn	Ser	Met	Ala	1/6
<i>p</i> -azidoPheRS-2	Val	Asn	Ser	Ala	Ala	1/6
p-azidoPheRS-3	Leu	Asn	Ser	Ala	Ala	1/6
<i>p</i> -azidoPheRS-4	Val	Asn	Ser	Ala	Val	1/6
<i>p</i> -azidoPheRS-5	Ile	Asp	Asn	Phe	Val	1/6
<i>p</i> -azidoPheRS-6	Thr	Asn	Ser	Ala	Leu	1/6

<sup>&</sup>lt;sup>a</sup> These clones also contain a Asp165Gly mutation

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# EXAMPLE 3: ADDING AMINO ACID WITH NOVEL REACTIVITY TO THE GENETIC CODE OF EUKARYOTES

[0320] A site-specific, fast, reliable, and irreversible method of bioconjugation to proteins based on a [3+2] cycloaddition is demonstrated. There is a considerable need for chemical reactions that modify proteins under physiological conditions in a highly selective fashion. See, e.g., Lemineux, & Bertozzi, (1996) TIBTECH, 16:506-513. Most reactions currently used for the selective modification of proteins involve covalent bond formation between nucleophilic and electrophilic reaction partners, e.g. the reaction of α-haloketones with histidine or cysteine side chains. Selectivity in these cases is determined by the number and accessibility of the nucleophilic residues in the protein. In the case of synthetic or semisynthetic proteins, other more selective reactions can be used such as the reaction of an unnatural keto-amino acid with hydrazides or aminooxy compounds. See, e.g., Cornish,

et al., (1996) Am. Chem. Soc., 118:8150-8151; and, Mahal, et al., (1997) Science, 276:1125-1128. Recently, it has been possible to genetically encode unnatural amino acids (see, e.g., Wang, et al., (2001) Science 292:498-500; Chin, et al., (2002) Am. Chem. Soc. 124:9026-9027; and, Chin, et al., (2002) Proc. Natl. Acad. Sci., 99:11020-11024), including ketone containing amino acids (see, e.g., Wang, et al., (2003) Proc. Natl. Acad. Sci., 100:56-61; Zhang, et al., (2003) Biochemistry, 42:6735-6746; and, Chin, et al., (2003) Science, in press), in bacteria and yeast using orthogonal tRNA-synthetase pairs with altered amino acid specificities. This methodology has made possible the selective labeling of virtually any protein with a host of reagents including fluorophores, crosslinking agents and cytotoxic molecules.

A highly efficient method for the selective modification of proteins is [0321] described, which involves the genetic incorporation of azide or acetylene containing unnatural amino acids into proteins in response to, e.g., the amber nonsense codon, TAG. These amino acid side chains can then be modified by a Huisgen [3+2] cycloaddition reaction (see, e.g., Padwa, A. in Comprehensive Organic Synthesis, Vol. 4, (1991) Ed. Trost, B. M., Pergamon, Oxford, p. 1069-1109; and, Huisgen, R. in 1,3-Dipolar Cycloaddition Chemistry, (1984) Ed. Padwa, A., Wiley, New York, p. 1-176) with alkynyl (acetylene) or azide derivatives, respectively. Because this method involves a cycloaddition rather than a nucleophilic substitution, proteins can be modified with extremely high selectivity (another method that can be used is the ligand exchange on a bisarsenic compound with a tetracysteine motif, see, e.g., Griffin, et al., (1998) Science 281:269-272). This reaction can be carried out at room temperature in aqueous conditions with excellent regioselectivity (1,4 > 1,5) by the addition of catalytic amounts of Cu(I) salts to the reaction mixture. See, e.g., Tornoe, et al., (2002) Org. Chem. 67:3057-3064; and, Rostovtsev, et al., (2002) Angew. Chem. Int. Ed. 41:2596-2599. Indeed, Finn and coworkers have shown that this azide-alkyne [3+2] cycloaddition can be conducted on the surface of an intact cowpea mosaic virus. See, e.g., Wang, et al., (2003) J. Am. Chem. Soc., 125:3192-3193. For another recent example of the electrophilic introduction of an azido group into a protein and a subsequent [3+2] cycloaddition, see, e.g., Speers, et al., (2003) J. Am. Chem. Soc., 125:4686-4687.

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[0322] In order to selectively introduce either the alkynyl (acetylene) or azide functional group into eukaryotic proteins at unique sites, evolved orthogonal TyrRS/tRNA<sub>CUA</sub> pairs were generated in yeast that genetically encode the acetylene and

azido amino acids, **Figure 11**, **1** and **2**, respectively. The resulting proteins can be efficiently and selectively labeled with fluorophores in a subsequent cycloaddition reaction under physiological conditions.

Previously, an E. coli tyrosyl tRNA-tRNA synthetase pair was demonstrated [0323] as being orthogonal in yeast, i.e., neither the tRNA nor the synthetase cross react with the 5 endogenous yeast tRNA or synthetases. See, e.g., Chin, et al., (2003) Chem. Biol., 10:511-519. This orthogonal tRNA-synthetase pair has been used to selectively and efficiently incorporate a number of unnatural amino acids in yeast in response to the TAG codon (e.g., Chin, et al., (2003) Science, in press). In order to alter the amino acid specificity of the E. coli tyrosyl-tRNA synthetase to accept amino acid 1 or 2 of Figure 11, a library of ~10<sup>7</sup> 10 mutants was generated by randomizing the codons for Tyr<sup>37</sup>, Asn<sup>126</sup>, Asp<sup>182</sup>, Phe<sup>183</sup>, and Leu<sup>186</sup>. These five residues were chosen based on a crystal structure of the homologous synthetase from B. stearothermophilus. To obtain a synthetase for which the particular amino acid serves as a substrate, a selection scheme was used in which the codons for Thr<sup>44</sup> and Arg<sup>110</sup> of the gene for the transcriptional activator GAL4 were converted to amber 15 nonsense codons (TAG). See, e.g., Chin, et al., (2003) Chem. Biol., 10:511-519. Suppression of these amber codons in the MaV203:pGADGAL4(2TAG) yeast strain leads to production of full length GAL4 (see, e.g., Keegan, et al., (1986) Science, 231:699-704; and, Ptashne, (1988) Nature, 335:683-689) which in turn drives expression of the HIS3 and URA3 reporter genes. The latter gene products complement histidine and uracil auxotrophy 20 allowing clones harboring active synthetase mutants to be selected in the presence of 1 or 2 of Figure 11. Synthetases that load endogenous amino acids are removed by growth on medium lacking 1 or 2 of Figure 11 but containing 5-fluoroorotic acid, which is converted into a toxic product by URA3. By passing the library through three rounds of selection (positive, negative, positive), we identified synthetases selective for 1 of Figure 11 (pPR-25 EcRS1-5) and for 2 of Figure 11 (pAZ-EcRS1-6) as shown in Table 8.

[0324] All synthetases show a strong sequence similarity, including a conserved  $Asn^{126}$ , suggesting an important functional role for this residue. Surprisingly, the synthetases pPR-EcRS-2 and pAZ-EcRS-6, evolved to bind 1 and 2 of Figure 11 respectively, converged to the same sequence  $(Tyr^{37} \rightarrow Thr^{37}, Asn^{126} \rightarrow Asn^{126}, Asp^{182} \rightarrow Ser^{182}$ , and  $Phe^{183} \rightarrow Ala^{183}$ ,  $Leu^{186} \rightarrow Leu^{186}$ ). Both hydrogen bonds between the phenolic hydroxy group of bound tyrosine and  $Tyr^{37}$  and  $Asp^{182}$  are disrupted by mutations to Thr and Ser, respectively.  $Phe^{183}$  is converted to Ala, possibly providing more space for the

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accommodation of the unnatural amino acid. To confirm the ability of this synthetase (and the other synthetases) to accept either amino acid as a substrate selection strains harboring the synthetase plasmids were grown on media lacking uracil (the same results were obtained for media lacking histidine) but supplemented with either 1 or 2 of Figure 11. Growth results revealed that four of the five alkyne synthetases were able to load both unnatural amino acids onto its tRNA. The azido synthetases seem to be more selective, since only pAZ-EcRS-6 (which is identical with pPR-EcRS-2) was able to amino acylate its tRNA with both 1 and 2 of Figure 11. The fact that no growth was detected in the absence of 1 or 2 of Figure 11 suggests that the synthetases do not accept any of the 20 common amino acids as a substrate. See Figure 14.

[0325] For all further experiments pPR-EcRS-2 (pAZ-EcRS-6) was used, allowing one to control which unnatural amino acid is incorporated simply by adding either 1 or 2 of Figure 11 to media containing the expression strain. For protein production the codon for the permissive residue Trp<sup>33</sup> of human superoxide dismutase-1 (SOD) fused to a C-terminal 6×His tag was mutated to TAG. For example, human superoxide dismutase (Trp<sup>33</sup> TAG) HIS was cloned between the GAL1 promoter and CYC1 terminator from pYES2.1 (Invitrogen, Carlsbad, CA USA). Mutant synthetase and tRNA genes on pECTyrRS-tRNA<sub>CUA</sub> derived plasmids were co-transformed with pYES2.1 SOD(Trp<sup>33</sup> TAG) HIS into the strain InvSc (Invitrogen). For protein expression, cells were grown in SD –trp, –ura +raffinose and expression was induced at an OD<sub>660</sub> of 0.5 by the addition of galactose. Protein was expressed in the presence or absence of 1 mM 1 or 2 of Figure 11 and purified by Ni-NTA chromatography (Qiagen, Valencia, CA, USA).

[0326] Analysis by SDS-PAGE and Western blot revealed unnatural amino acid dependent protein expression with a fidelity of >99% as judged by densitometry comparisons to protein expression in absence of 1 or 2 of Figure 11. See Figure 12. To further confirm the identity of the amino acid incorporated, a tryptic digest was subjected to liquid chromatography and tandem mass spectrometry.

[0327] For example, the wild-type and mutant hSOD were purified using nickel affinity column and protein bands were visualized by colloidal Coomassie stain. Gels bands corresponding to wild-type and mutant SOD were excised from polyacrylamide gels, sliced into 1.5-mm cubes, reduced and alkylated, then subjected to trypsin hydrolysis essentially as described. See, e.g., Shevchenko, A et al., (1996) Anal. Chem. 68:850-858. Tryptic

peptides containing the unnatural amino acid were analyzed by nanoflow reversed-phase HPLC/µESI/MS with an LCQ ion trap mass spectrometer. See, Figure 15, Panel A and B. Liquid chromatography tandem mass spectrometry (LC-MS/MS) analysis was performed on a Finnigan LCQ Deca ion trap mass spectrometer (Thermo Finnigan) fitted with a Nanospray HPLC (Agilent 1100 series).

[0328] The precursor ions corresponding to the singly and doubly charged precursor ions of the peptide VY\*GSIK (SEQ ID NO:87) containing the unnatural amino acid (denoted Y\*) were separated and fragmented with an ion trap mass spectrometer. The fragment ion masses could be unambiguously assigned, confirming the site-specific incorporation of each unnatural amino acid. LC MS/MS did not indicate incorporation of any natural amino acid at this position. The signal-to-noise of the peptide for all mutants were >1000 suggesting fidelity of incorporation better than 99.8%. See, Figure 15, Panel A and B.

[0329] To demonstrate that small organic molecules can be conjugated to proteins by an azide-alkyne [3+2] cycloaddition reaction, the dyes 3-6 indicated in Figure 13, Panel A, which contain either an acetylenic or an azido group and bear a dansyl or fluoresceine fluorophore, were synthesized (see Example 5 herein). The cycloaddition itself was carried out with 0.01 mM protein in phosphate buffer (PB), pH 8, in the presence of 2 mM 3-6 indicated in Figure 13, Panel A, 1 mM CuSO<sub>4</sub>, and ~1 mg Cu-wire for 4 hours at 37 °C (see Figure 13, Panel B).

[0330] For example, to 45 μL of protein in PB buffer (pH = 8) was added 1 μL of CuSO<sub>4</sub> (50 mM in H<sub>2</sub>O), 2 μL of dye (50 mM in EtOH), 2 μL of tris(1-benzyl-1*H*-[1,2,3]triazol-4-ylmethyl)amine (50 mM in DMSO), and Cu wire. After 4 hours at room temperature or 37 °C or overnight at 4 °C, 450 μL H<sub>2</sub>O were added and the mixture was spun through a dialysis membrane (10 kDa cut off). After washing the supernatant with 2×500 μL by centrifugation, the solution was brought to a volume of 50 mL. A sample of 20 mL was analyzed by SDS-PAGE. Occasionally remaining dye could be removed from the gel by soaking in H<sub>2</sub>O/MeOH/AcOH (5:5:1) overnight. The use of tris(carboxyethyl)phosphine as the reducing agent generally led to less efficient labeling. In contrast to earlier observations (e.g., Wang, Q. et al., (2003) J. Am. Chem. Soc. 125:3192-3193), the presence or absence of the tris(triazolyl)amine ligand did not have a substantial influence on the outcome of the reaction.

[0331] After dialysis the labeled proteins were then analyzed by SDS-PAGE and ingel imaged using a densitometer in case of the dansyl dyes 3-4 indicated in Figure 13,
Panel A (λ<sub>ex</sub> = 337 nm, λ<sub>em</sub> = 506 nm) or a phosphorimager in the case of the fluoresceine dyes 5-6 indicated in Figure 13, Panel A (λ<sub>ex</sub> = 483 nm, λ<sub>em</sub> = 516 nm). See, e.g., Blake,
(2001) Curr. Opin. Pharmacol., 1:533-539; Wouters, et al., (2001) Trends in Cell Biology 11:203-211; and, Zacharias, et al., (2000) Curr. Opin. Neurobiol., 10:416-421. The labeled proteins were characterized by LC MS/MS analysis of tryptic digests showing site-specific attachment of the fluorophores and the conversion was 75% on average (e.g., as determined by comparison of A<sub>280</sub>/A<sub>495</sub> values for SOD labeled with 5 or 6 indicated in Figure 13,
Panel A). The selectivity of this bioconjugation is verified by the fact that there was no observable reaction between 3 indicated in Figure 13, Panel A and alkyne protein or 4 indicated in Figure 13, Panel A and azido protein.

TABLE 8 EVOLVED SYNTHETASES.

pPR-EcRS selected for 1 and pAZ-EcRS selected for 2 (as indicated in Figure 11)

synthetase	37	126	182	183	186
wild type	Tyr	Asn	Asp	Phe	Leu
pPR-EcRS-1	Gly	Asn	Ser	Met	Leu
pPR-EcRS-2	Thr	Asn	Ser	Ala	Leu
pPR-EcRS-3	Ser	Asn	Thr	Met	Val
pPR-EcRS-4	Ala	Asn	Ser	Tyr	Leu
pPR-EcRS-5	Ala	Asn	Thr	Met	Cys
pAZ-EcRS-1	Leu	Asn	Ser	Met	Ala
pAZ-EcRS-2	Val	Asn	Ser	Ala	Ala
pAZ-EcRS-3	Leu	Asn	Ser	Ala	Ala
pAZ-EcRS-4	Val	Asn	Ser	Ala	Val
pAZ-EcRS-5	Ile	Asp	Asn	Phe	Val
pAZ-EcRS-6	Thr	Asn	Ser	Ala	Leu

### 15 EXAMPLE 4: SYNTHESIS OF AN ALKYNE AMINO ACID

[0332] In one aspect of the invention, the invention provides alkynyl amino acids. An example of a structure of the alkynyl amino acid is illustrated by Formula IV:

$$R_2$$
— $R_1$ — $CO_2H$ 
 $NH_2$ 

[0333] An alkyne amino acid is typically any structure having Formula IV, where  $R_1$  is a substituent used in one of the twenty natural amino acids and  $R_2$  is an alkynyl substituent. For example, 1 in Figure 11 illustrates the structure of *para*-propargyloxyphenylalanine. *p*-Propargyloxyphenylalanine can be synthesized, e.g., as outline below. In this embodiment, the synthesis of *p*-propargyloxyphenylalanine can be

completed in three steps starting from the commercially available N-Boc-tyrosine.

[0334] For example, N-tert-butoxycarbonyl-tyrosine (2 g, 7 mmol, 1 equiv.) and  $K_2CO_3$  (3 g, 21 mmol, 3 equiv.) were suspended in anhydrous DMF (15 mL). Propargyl bromide (2.1 mL, 21 mmol, 3 equiv., 80% solution in toluene) was slowly added and the reaction mixture was stirred for 18 hours at room temperature. Water (75 mL) and  $Et_2O$  (50 mL) were added, the layers were separated and the aqueous phase was extracted with  $Et_2O$  (2 × 50 mL). The combined organic layers were dried (MgSO<sub>4</sub>) and the solvent was removed under reduced pressure. The product was obtained as a yellow oil (2.3 g, 91%) and used in the next step without further purification. The Boc-protected product is illustrated

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below as chemical structure 8:

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2-tert-Butoxycarbonylamino-3-[4-(prop-2-ynyloxy)phenyl]-propionic acid propargyl ester

20 [0335] Acetyl chloride (7 mL) was added carefully to methanol (60 mL) at 0 °C to give a 5 M solution of anhydrous HCl in MeOH. The product of the previous step (2 g, 5.6 mmol) was added and the reaction was stirred for 4 hours while it was allowed to warm to ambient temperature. After removing the volatiles under reduced pressure, a yellowish solid (1.6 g, 98%) (see chemical structure 9) was obtained which was directly used in the next step.

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### 2-Amino-3-[4-(prop-2-ynyloxy)phenyl]-propionic acid propargyl ester

[0336] The propargyl ester (1.6 g, 5.5 mmol) from the previous step was dissolved in a mixture of aqueous 2 N NaOH (14 mL) and MeOH (10 mL). After stirring for 1.5 h at room temperature, the pH was adjusted to 7 by adding conc. HCl. Water (20 mL) was added and the mixture was kept at 4 °C overnight. The precipitate was filtered, washed with ice-cold H<sub>2</sub>O, and dried under vacuum yielding, 1.23 g (90%) of 1 in **Figure 11** (2-Amino-3-phenylpropionic acid (1) (also known as *p*-propargyloxyphenylalanine) as a white solid.  $^{1}$ H NMR (400 MHz, D<sub>2</sub>O) (as the potassium salt in D<sub>2</sub>O)  $\delta$  7.20 (d, J = 8.8 Hz, 2 H), 6.99 (d, J = 8.8 Hz, 2 H), 4.75 (s, 2 H), 3.50 (dd, J = 5.6, 7.2 Hz, 1 H), 2.95 (dd, J = 5.6, 13.6 Hz, 1 H), 2.82 (dd, J = 7.2, 13.6 Hz, 1 H);  $^{13}$ C NMR (100 MHz, D<sub>2</sub>O)  $\delta$  181.3, 164.9, 155.6, 131.4, 130.7, 115.3, 57.3, 56.1, 39.3; HRMS (CI) m/z 220.0969 [C<sub>12</sub>H<sub>13</sub>NO<sub>3</sub> (M+1) requires 220.0968].

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# EXAMPLE 5: ADDITION OF MOLECULES TO PROTEINS WITH AN UNNATURAL AMINO ACID THROUGH A [3+2] CYCLOADDITION

[0337] In one aspect, the invention provides methods and related compositions of proteins comprising unnatural amino acids coupled to additional substituent molecules. For example, additional substituents can be added to an unnatural amino acid through a [3+2] cycloaddition. See, e.g., Figure 16. For example, the [3+2] cycloaddition of a desired molecule (e.g., that include a second reactive group, such as an alkyne triple bond or azido group) to a protein with an unnatural amino acid (e.g., having a first reactive group, such as azido group or triple bond) can be done following published conditions for the [3+2] cycloaddition reaction. For example, a protein comprising the unnatural amino acid in PB buffer (pH = 8) is added to CuSO<sub>4</sub>, the desired molecule, and Cu wire. After the mixture is incubated (e.g., about 4 hours at room temperature or 37 °C, or overnight at 4 °C), H<sub>2</sub>O is added and the mixture is filtered through a dialysis membrane. The sample can be analyzed for the addition, e.g., by gel analysis.

[0338] Examples of such molecules include, but are not limited to, e.g., a molecule having a triple bond or azido group, such as molecules have the structure of Formula 3, 4, 5, and 6 of Figure 13, Panel A and the like. Furthermore, triple bonds or azido groups can be incorporated into the structures of other molecules of interest, such as polymers (e.g., poly(ethylene glycol) and derivatives), crosslinking agents, additional dyes, photocrosslinkers, cytotoxic compounds, affinity labesl, biotin, saccharides, resins, beads, a

second protein or polypeptide, metal chelators, cofactors, fatty acids, carbohydrates, polynucleotides (e.g., DNA, RNA, etc.), and the like, which then can also be used in [3+2] cycloadditions.

In one aspect of the invention, molecules having the Formula 3, 4, 5, or 6 of Figure 13, Panel A can be synthesized as described below. For example, an alkyne dye as shown in 3 of Figure 13, Panel A and in chemical structure 3 below was synthesized by adding propargylamine (250 μL, 3.71 mmol, 3 equiv.) to a solution of dansyl chloride (500 mg, 1.85 mmol, 1 equiv.) and triethylamine (258 μL, 1.85 mmol, 1 equiv.) in CH<sub>2</sub>Cl<sub>2</sub> (10 mL) at 0 °C. After stirring for 1 hour, the reaction mixture was warmed to room temperature and stirred for an additional hour. The volatiles were removed *in vacuo* and the crude product was purified by chromatography on silica gel (Et<sub>2</sub>O/hexanes = 1:1) yielding 3 of Figure 13, Panel A (418 mg, 78%) as a yellow solid. The analytical data are identical with those reported in the literature. See, for example, Bolletta, F et al., (1996) Organometallics 15:2415-17. An example of a structure of an alkyne dye that can be used in the invention is shown in chemical structure 3:

[0340] An azido dye as shown as shown in 4 of Figure 13, Panel A and in chemical structure 4 below was synthesized by adding 3-azidopropylamine (e.g., as described by Carboni, B et al., (1993) J. Org. Chem. 58:3736-3741) (371 mg, 3.71 mmol, 3 equiv.) to a solution of dansyl chloride (500 mg, 1.85 mmol, 1 equiv.) and triethylamine (258  $\mu$ L, 1.85 mmol, 1 equiv.) in CH<sub>2</sub>Cl<sub>2</sub> (10 mL) at 0 °C. After stirring for 1 hour, the reaction mixture was warmed to room temperature and stirred for an additional hour. The volatiles were removed in vacuo and the crude product was purified by chromatography on silica gel (Et<sub>2</sub>O/hexanes = 1:1) yielding 4 of Figure 13, Panel A (548 mg, 89%) as a yellow oil. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  8.55 (d, J = 8.4 Hz, 1 H), 8.29 (d, J = 8.8 Hz, 1 H), 8.23 (dd, J = 1.2, 7.2 Hz, 1 H), 7.56-7.49 (comp, 2 H), 7.18 (d, J = 7.6 Hz, 1 H), 5.24 (br s, 1 H), 3.21 (t, J = 6.4 Hz, 2 H), 2.95 (dt, J = 6.4 Hz, 2 H), 2.89 (s, 6 H), 1.62 (quin, J = 6.4 Hz, 2 H);

<sup>13</sup>C NMR (100 MHz, CDCl<sub>3</sub>) δ 134.3, 130.4, 129.7, 129.4, 128.4, 123.3, 118.8, 115.3, 48.6, 45.4, 40.6, 28.7 (not all signals of quaternary carbon atoms are visible in the <sup>13</sup>C NMR spectrum); HRMS (CI) m/z 334.1336 [C<sub>15</sub>H<sub>20</sub>N<sub>5</sub>O<sub>2</sub>S (M+1) requires 334.1332]. An example of a structure of an azido dye is shown in chemical structure 4:

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[0341] An alkyne dye as shown in 5 of Figure 13, Panel A and in chemical 10 structure 5 below was synthesized by adding EDCI (1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride) (83 mg, 0.43 mmol, 1 equiv.) to a solution of fluoresceinamine (150 mg, 0.43 mmol, 1 equiv.) and 10-undecynoic acid (79 mg, 0.43, 1 equiv.) in pyridine (2 mL) at room temperature. The suspension was stirred overnight and the reaction mixture was poured into  $H_2O$  (15 mL). The solution was acidified (pH < 2) by adding conc. HCl. 15 After stirring for 1 h, the precipitate was filtered off, washed with H<sub>2</sub>O (5 mL) and dissolved in small amount of EtOAc. Addition of hexanes led to the precipitation of 5 of Figure 13, Panel A as orange crystals, which were collected and dried under vacuum (138 mg, 63%). The analytical data are identical with those reported in the literature. See, e.g., Crisp, G. T.; & Gore, J. (1997) Tetrahedron 53:1505-1522. An example of a structure of an alkyne dye is shown in chemical structure 5: 20

25 [0342] An azido dye as shown in 6 of Figure 13, Panel A and in chemical structure 6 below was synthesized by adding EDCI (83 mg, 0.43 mmol, 1 equiv.) to a solution of fluoresceinamine (150 mg, 0.43 mmol, 1 equiv.) and 4-(3-azidopropylcarbamoyl)-butyric

acid (e.g., synthesized by reacting 3-azidopropylamine with glutaric acid anhydride) (92 mg, 0.43, 1 equiv.) in pyridine (2 mL) at room temperature. The suspension was stirred over night and the reaction mixture was poured in  $H_2O$  (15 mL). The solution was acidified (pH < 2) by adding conc. HCl. After stirring for 1 hour, the precipitate was filtered off, washed with 1 N HCl (3×3 mL) and was dissolved in a small amount of EtOAc. Addition of hexanes led to the precipitation of **6** of **Figure 13**, **Panel A** as orange crystals, which were collected and dried under vacuum (200 mg, 86%). <sup>1</sup>H NMR (400 MHz, CD<sub>3</sub>OD)  $\delta$  8.65 (s, 1 H), 8.15 (d, J = 8.4 Hz, 1 H), 7.61-7.51 (comp, 2 H), 7.40 (d, J = 8.4 Hz, 1 H), 7.35 (br s, 2 H), 7.22-7.14 (comp, 2 H), 6.85-6.56 (comp, 3 H), 3.40-3.24 (comp, 4 H), 2.54 (t, J = 7.2 Hz, 2 H), 2.39-2.30 (comp, 2 H), 2.10-1.99 (comp, 2 H), 1.82-1.72 (comp, 2 H); <sup>13</sup>C NMR (100 MHz, CD<sub>3</sub>OD)  $\delta$  175.7, 174.4, 172.4, 167.9, 160.8, 143.0, 134.3, 132.9, 131.8, 129.6, 124.4, 123.3, 121.1, 118.5 103.5, 50.2, 38.0, 37.2, 36.2, 29.8, 22.9 (not all signals of quaternary carbon atoms are visible in the <sup>13</sup>C NMR spectrum ); HRMS (CI) m/z 544.1835 [C<sub>28</sub>H<sub>25</sub>N<sub>5</sub>O<sub>7</sub> (M+1) requires 544.1827]. An example of a structure of an azido dye is shown in chemical structure 6:

[0343] In one embodiment, a PEG molecule can also be added to a protein with an unnatural amino acid, e.g., an azido amino acid or a propargyl amino acid. For example, a propargyl amide PEG (e.g., illustrated in **Figure 17**, **Panel A**) can be added to a protein with an azido amino acid through a [3+2] cycloaddition. See e.g., **Figure 17**, **Panel A**. **Figure 17**, **Panel B** illustrates a gel analysis of a protein with an added PEG substituent.

[0344] In one aspect of the invention, a propargyl amide PEG (e.g., illustrated in Figure 17, Panel A) can be synthesized as described below. For example, a solution of propargylamine (30 μL) in CH<sub>2</sub>Cl<sub>2</sub> (1 mL) was added to the 20 kDa PEG-hydroxysuccinimide ester (120 mg, purchased from Nektar). The reaction was stirred for 4 hours at room temperature. Then Et<sub>2</sub>O (10 mL) was added, the precipitate was filtered off,

and was twice recrystallized from MeOH (1 mL) by addition of Et<sub>2</sub>O (10 mL). The product was dried under vacuum furnishing a white solid (105 mg, 88% yield). See, e.g., Figure 17, Panel C.

### EXAMPLE 6: EXEMPLARY O-RSs AND O-tRNAs.

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[0345] An exemplary O-tRNA comprises SEQ ID NO.:65 (See, Table 5). Example O-RSs include SEQ ID NOs.: 36-63, 86 (See, Table 5). Examples of polynucleotides that encode O-RSs or portions thereof (e.g., the active site) include SEQ ID NOs.: 3-35. In addition, exemplary amino acid changes of O-RSs are indicated in Table 6.

Table 6: Evolved EcTyrRS Variants

Residue #	37	126	182	183	186	Representation
Ec TyrRS	Tyr	Asn	Asp	Phe	Leu	
n indeDhaDC 1	Val	Aon	Cor	T. #	Lou	1/8
p-iodoPheRS-1	Val	Asn	Ser	Tyr	Leu	
p-iodoPheRS-2	lle	Asn	Ser	Met	Leu	1/8
p-iodoPheRS-3	Val	Asn	Ser	Met	Ala	6/8
OMeTyrRS-1	Val	Asn	Ser	Met	Leu	5/13
OMeTyrRS-2	Thr	Asn	Thr	Met	Leu	1/13
OMeTyrRS-3	Thr	Asn	Thr	Tyr	Leu	1/13
OMeTyrRS-4	Leu	Asn	Ser	Met	Ser	1/13
OMeTyrRS-5	Leu	Asn	Ser	Met	Ala	1/13
OMeTyrRS-6	Thr	Asn	Arg	Met	Leu	4/13
J. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	••••		79			
p-acetylPheRS-1	lle	Asn	Gly	Met	Ala	4/4
p-acetylPheRS-1 <sup>a</sup>	lle	Asn	Gly	Met	Ala	10/10
,						
p-benzoylPheRS-1	Gly	Asn	Gly	Phe	Ala	1/2
p-benzoylPheRS-2	Gly	Asn	Gly	Tyr	Met	1/2
p-azidoPheRS-1	Leu	Asn	Ser	Met	Ala	1/6
p-azidoPheRS-2	Val	Asn	Ser	Ala	Ala	1/6
p-azidoPheRS-3	Leu	Asn	Ser	Ala	Ala	1/6
p-azidoPheRS-4	Val	Asn	Ser	Ala	Val	1/6
p-azidoPheRS-5	lle	Asp	Asn	Phe	Val	1/6
p-azidoPheRS-6	Thr	Asn	Ser	Ala	Leu	1/6
			_			
p-PR-EcRS-1	Gly	Asn	Ser	Met	Leu	1/10
p-PR-EcRS-2	Thr	Asn	Ser	Ala	Leu	1/10
p-PR-EcRS-3	Ser	Asn	Thr	Met	Val	1/10
p-PR-EcRS-4	Ala	Asn	Ser	Tyr	Leu	1/10
p-PR-EcRS-5	Ala	Asn	Thr	Met	Cys	1/10
p-PR-EcRS-6	Thr	Asn	Thr	Phe	Met	1/10
p-PR-EcRS-7	Thr	Asn	Ser	Val	Leu	1/10
p-PR-EcRS-8	Val	Asn	Ser	Met	Thr	1/10
p-PR-EcRS-9	Ser	Asn	Ser	Phe	Leu	1/10
p-PR-EcRS-10	Thr	Asn	Thr	Phe	Thr	1/10

<sup>&</sup>lt;sup>a</sup> These clones also contain a Asp165Gly mutation

- [0346] It is understood that the examples and embodiments described herein are for illustrative purposes only and that various modifications or changes in light thereof will be suggested to persons skilled in the art and are to be included within the spirit and purview of this application and scope of the appended claims.
- 5 [0347] While the foregoing invention has been described in some detail for purposes of clarity and understanding, it will be clear to one skilled in the art from a reading of this disclosure that various changes in form and detail can be made without departing from the true scope of the invention. For example, all the techniques and apparatus described herein can be used in various combinations. All publications, patents, patent applications, and/or other documents cited in this application are incorporated by reference in their entirety for all purposes to the same extent as if each individual publication, patent, patent application, and/or other document were individually indicated to be incorporated by reference for all purposes.

TABLE 5

SEQ ID NO.:	Label	SEQUENCE
SEQ ID	E. coli wild-	ATGGCAAGCAGTAACTTGATTAAACAATTGCAAGAGCGGGGGCTGGTA
NO.: 1	type TyrRS	GCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGACTGGCGCAAGG
}	(synthetase)	CCCGATCGCGCTCTATTGCGGCTTCGATCCTACCGCTGACAGCTTGCAT
	polynucleotide	TTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGGCGG
		GCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCG
		ACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTG
		TTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCG
		ATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCGAACAACTATGACT
		GGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACA
		CTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCT
}		CAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAACCTG
i		TTGCAGGGTTATGACTTCGCCTGTCTGAACAAACAGTACGGTGTGGTGC
İ		TGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTCTGGTATCGA
		CCTGACCCGTCGTCTGCATCAGAATCAGGTGTTTTGGCCTGACCGTTCCG
1		CTGATCACTAAAGCAGATGGCACCAAATTTGGTAAAACTGAAGGCGGC
		GCAGTCTGGTTGGATCCGAAGAAACCAGCCCGTACAAATTCTACCAG
		TTCTGGATCAACACTGCGGATGCCGACGTTTACCGCTTCCTGAAGTTCT
		TCACCTTTATGAGCATTGAAGAGATCAACGCCCTGGAAGAAGAAGATA
		AAAACAGCGGTAAAGCACCGCGCGCCCAGTATGTACTGGCGGAGCAG
		GTGACTCGTCTGGTTCACGGTGAAGAAGGTTTACAGGCGGCAAAACGT
		ATTACCGAATGCCTGTTCAGCGGTTCTTTGAGTGCGCTGAGTGAAGCGG
		ACTTCGAACAGCTGGCGCAGGACGGCGTACCGATGGTTGAGATGGAAA
		AGGGCGCAGACCTGATGCAGGCACTGGTCGATTCTGAACTGCAACCTT
		CCCGTGGTCAGGCACGTAAAACTATCGCCTCCAATGCCATCACCATTAA
}		CGGTGAAAAACAGTCCGATCCTGAATACTTCTTTAAAGAAGAAGATCG
	:	TCTGTTTGGTCGTTTTACCTTACTGCGTCGCGGTAAAAAGAATTACTGT
I		CTGATTTGCTGGAAATAA
SEQ ID	E. coli wild-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALYCGFDPTADSLHLGH
NO.: 2	type TyrRS	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
110 2	(synthetase)	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYDFACLNKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	(aa)	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pOMe-1	ATGGCAAGCAGTAACTTGATTAAACAATTGCAAGAGCGGGGGCTGGTA
NO.: 3	Synthetase	GCCCAGGTGACGAGGGAAGCGTTAGCAGAGCGACTGGCGCAAGGC
140 3	1	CCGATCGCACTCGTGTGGCTTCGATCCTACCGCTGACAGCTTGCATTT
	polynucleotide	GGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGGCGGGC
		CACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGAC
		CCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTT
		CAGGAGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATT
		TCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATAATTATGACTGGTT
		CGGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTC
		TCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAAC
	i	CGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGC
	1	AGGGTTATAGTATGGCCTGTTTGAACAAACAGTACGGTGTGGTGCTGCA
		AATTGGTGGTTCTGACCAGTGGGGTAACATCACTTCTGGTATCGACCTG
	1	ACCCGTCGTCTGCATCAGAATCAGTGTTTTGGCCTGACCGTTCCGCTGA
	I	TCACTAAAGCAGATGGCACCAAATTTGGTAAAACTGAAGGCGGCGCAG

		·
		TCTGGTTGGATCCGAAGAAAACCAGCCCGTACAAATTCTACCAGTTCTG GATCAACACTGCGGATGCCGACGTTTACCGCTTCCTGAAGTTCTTCACC TTTATGAGCATTGAAGAGATCAACGCCCTGGAAGAAGAAGATAAAAAC
		AGCGGTAAAGCACCGCGCGCCCAGTATGTACTGGCGGAGCAGGTGACT
		GAATGCCTGTTCAGCGGTTCTTTGAGTGCGCTGAGTGAAGCGGACTTCG
		AACAGCTGGCGCAGGACGGCGTACCGATGGTTGAGATGGAAAAGGGCG
		CAGACCTGATGCAGGCACTGGTCGATTCTGAACTGCAACCTTCCcGTGG
		TCAGGCACGTAAAACTATCGCCTCCAATGCCATCACCATTAACGGTGAA
		AAACAGTCCGATCCTGAATACTTCTTTAAAGAAGAAGATCGTCTGTTTG GTCGTTTTACCTTACTGCGTCGCGGTAAAAAGAATTACTGTCTGATTTGC
	-	TGGAAATAA
SEQ ID	pOMe-2	ATGGCAAGCAGTAACTTGATTAAACAATTGCAAGAGCGGGGGCTGGTA
NO.: 4	Synthetase	gCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGACTGGCGCAAGGC
	polynucleotide	CCGATCGCACTCACTTGTGGCTTCGATCCTACCGCTGACAGCTTGCATTT GGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGCGGGC
		CACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGAC
		CCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTT
		CAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATT
		TCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATAATTATGACTGGTT
		CAGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTC
		TCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAAC
		CGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGC AGGGTTATACGTATGCCTGTCTGAACAAACAGTACGGTGTGGTGCTGCA
		AATTGGTGGTTCTGACCAGTGGGGTAACATCACTTCTGGTATCGACCTG
		ACCCGTCGTCTGCATCAGAATCAGGTGTTTTGGCCTGACCGTTCCGCTGA
		TCACTAAAGCAGATGGCACCAAATTTGGTAAAACTGAAGGCGGCGCAG
		TCTGGTTGGATCCGAAGAAACCAGCCCGTACAAATTCTACCAGTTCTG
		GATCAACACTGCGGATGCCGACGTTTACCGCTTCCTGAAGTTCTTCACC
		TTTATGAGCATTGAAGAGATCAACGCCCTGGAAGAAGAAGATAAAAAC AGCGGTAAAGCACCGCGCGCCCCAGTATGTACTGGCGGAGCAGGTGACT
		CGTCTGGTTCACGGTGAAGAAGGTTTACAGGCGGCAAAACGTATTACC
		GAATGCCTGTTCAGCGGTTCTTTGAGTGCGCTGAGTGAAGCGGACTTCG
		AACAGCTGGCGCAGGACGGCGTACCGATGGTTGAGATGGAAAAGGGCG
		CAGACCTGATGCAGGCACTGGTCGATTCTGAACTGCAACCTTCCCGTGG
		TCAGGCACGTAAAACTATCGCCTCCAATGCCATCACCATTAACGGTGAA
		AAACAGTCCGATCCTGAATACTTCTTTAAAGAAGAAGATCGTCTGTTTG GTCGTTTTACCTTACTGCGTCGCGGTAAAAAGAATTACTGTCTGATTTGC
		TGGAAATAA
SEQ ID	pOMe-3	ATGGCAAGCAGTAACTTGATTAAACAATTGCAAGAGCGGGGGCTGGTA
NO.: 5	Synthetase	GCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGACTGGCGCAAGGC
	polynucleotide	CCGATCGCACTCGTGTGTGGCTTCGATCCTACCGCTGACAGCTTGCATTT
		GGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGGCGGGC CACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGAC
		CCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTT
		CAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATT
		TCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATAATTATGACTGGTT
		CGGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTC
		TCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAAC
		CGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGC
		AGGGTTATAGTATGGCCTGTTTGAACAAACAGTACGGTGTGGTGCTGCA AATTGGTGGTTCTGACCAGTGGGGTAACATCACTTCTGGTATCGACCTG
		ACCCGTCGTCTGCATCAGAATCAGGTGTTTGGCCTGACCGTTCCGCTGA
		TCACTAAAGCAGATGCACCAAATTTGGTAAAACTGAAGGCGGCGCAG
		TCTGGTTGGATCCGAAGAAACCAGCCCGTACAAATTCTACCAGTTCTG
		GATCAACACTGCGGATGCCGACGTTTACCGCTTCCTGAAGTTCTTCACC
1		TTTATGAGCATTGAAGAGATCAACGCCCTGGAAGAAGAAGATAAAAAC
		AGCGGTAAAGCACCGCGCGCCCAGTATGTACTGGCGGAGCAGGTGACT CGTCTGGTTCACGGTGAAGAAGGTTTACAGGCGGCAAAACGTATTACC
L	L	COTCTOOTTCACOOTOAAOAAOOTTTACAOOCOOCAAAACOTATTACC

		GAATGCCTGTTCAGCGGTTCTTTGAGTGCGCTGAGTGAAGCGGACTTCG AACAGCTGGCGCAGGACGGCGTACCGATGGTTGAGATGGAAAAGGGCG CAGACCTGATGCAGGCACTGGTCGATTCTGAACTGCAACCTTCCCGTGG TCAGGCACGTAAAACTATCGCCTCCAATGCCATCACCATTAACGGTGAA AAACAGTCCGATCCTGAATACTTCTTTAAAGAAGAAGATCGTCTGTTTG GTCGTTTTACCTTACTGCGTCGCGGTAAAAAGAATTACTGTCTGATTTGC TGGAAATAA
SEQ ID NO.: 6	pOMe-4 Synthetase polynucleotide	ATGGCAAGCAGTAACTTGATTAAACAATTGCAAGAgCGGGGGCTGGTA GCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGACTGGCGCAAGGC CCGATCGCACTCGTGTGTGGCTTCGATCCTACCGCTGACAGCTTGCATTT GGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGGCGGC CACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGAC CCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTT
		CAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATT TCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATAATTATGACTGGTT CGGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTC TCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAAC CGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGC AGGGTTATAGTATGGCCTGTTTGAACAAACAGTACGGTGTGGTGCTGCA
		AATTGGTGGTTCTGACCAGTGGGGTAACATCACTTCTGGTATCGACCTG ACCCGTCGTCTGCATCAGAATCAGGTGTTTTGGCCTGACCGTTCCGCTGA TCACTAAAGCAGATGGCACCAAATTTGGTAAAACTGAAGGCGGCGCAG TCTGGTTGGATCCGAAGAAAACCAGCCCGTACAAATTCTACCAGTTCTG GATCAACACTGCGGATGCCGACGTTTACCGCTTCCTGAAGTTCTTCACC TTTATGAGCATTGAAGAGATCAACGCCCTGGAAGAAGAAGATAAAAAC
		AGCGGTAAAGCACCGCGCGCCCAGTATGTACTGGCGGAGCAGGTGACT CGTCTGGTTCACGGTGAAGAAGGTTTACAGGCGGCAAAACGTATTACC GAATGCCTGTTCAGCGGTTCTTTGAGTGCGCTGAGTGAAgCGGACTTCG AACAGCTGGCGCAGGACGGCGTACCGATGGTTGAGATGGAAAAGGGCG CAGACCTGATGCAGGCACTGGTCGATTCTGAACTGCAACCTTCCCGTGG
SEQ ID	pOMe-5	TCAGGCACGTAAAACTATCGCCTCCAATGCCATCACCATTAACGGTGAA AAACAGTCCGATCCTGAATACTTCTTTAAAGAAGAAGATCGTCTGTTTG GTCGTTTTACCTTACTGCGTCGCGGTAAAAAGAATTACTGTCTGATTTGC TGGAAATAA ATGGCAAGCAGTAACTTGATTAAACAATTGCAAGAGCGGGGGCTGGTA
NO.: 7	Synthetase polynucleotide	gCCCAGGTGACGAGGAGGAGCGTTAGCAAGAGCGGGGCTGGTA gCCCAGGTGACGGACGAGGAGCGTTAGCAGAGCGACTGGCGCAAGGC CCGATCGCACTCACGTGTGGCTTCGATCCTACCGCTGACAGCTTGCATT TGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGGCGGG CCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGA CCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGT
		TCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGAT TTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATAATTATGACTGGT TCGGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACACTT CTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAA CCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAGCCTGCTG
		CAGGGTTATACGATGGCCTGTCTGAACAAACAGTACGGTGTGGTGCTGC AAATTGGTGGTCTGACCAGTGGGGTAACATCACTTCTGGTATCGACCT GACCCGTCGTCTGCATCAGAATCAGGTGTTTGGCCTGACCGTTCCGCTG ATCACTAAAGCAGATGGCACCAAATTTGGTAAAACTGAAGGCGGCGCA GTCTGGTTGGATCCGAAGAAAACCAGCCCGTACAAATTCTACCAGTTCT
		GGATCAACACTGCGGATGCCGACGTTTACCGCTTCCTGAAGTTCTTCAC CTTTATGAGCATTGAAGAGATCAACGCCCTGGAAGAAGAAGATAAAAA CAGCGGTAAAGCACCGCGCGCCCAGTATGTACTGGCGGAGCAGGTGAC TCGTCTGGTTCACGGTGAAGAAGGTTTACAGGCGGCAAAACGTATTACC GAATGCCTGTTCAGCGGTTCTTTGAGTGCGCTGAGTGAAGCGGACTTCG
		AACAGCTGGCGCAGGACGGCGTACCGATGGTTGAGATGGAAAAGGGCG CAGACCTGATGCAGGCACTGGTCGATTCTGAACTGCAACCTTCCCGTGG TCAGGCACGTAAAACTATCGCCTCCAATGCCATCACCATTAACGGTGAA

		A A A A A A A A A A A A A A A A A A A
		AAACAGTCCGATCCTGAATACTTCTTTAAAGAAGAAGATCGTCTGTTTG GTCGTTTTACCTTACTGCGTCGCGGTAAAAAGAATTACTGTCTGATTTGC TGGAAATAA
SEQ ID NO.: 8	pOMe-6 (active site) Synthetase polynucleotide	CGGGGGCTGGTAGCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGA CTGGCGCAAGGCCCGATCGCACTCACTTGTGGCTTCGATCCTACCGCTG ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA ATTATGACTGGTTCAGCAATATGAATGTGCTGACCTTCCTGCGCGATAT TGGCAAACACTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGATCAGCGATTTCCC TACAACCTGCTGCAGGGTTATACCTATGCTTCACTGAGTTTTCC GTGTG
SEQ ID	pOMe-7	CGGGGGCTGGTACCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
NO.: 9	(active site) Synthetase polynucleotide	CTGGCGCAAGGCCCGATCGCACTCACTTGTGGCTTCGATCCTACCGCTG ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA ATTATGACTGGTTCAGCAATATGAATGTGCTGACCTTCCTGCGCGATAT TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC TACAACCTGCTGCAGGGTTATACGTATGCCTGTCTGAACAAACA
SEQ ID NO.: 10	pOMe-8 (active site) Synthetase polynucleotide	CGGGGGCTGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA CTGGCGCAAGGCCCGATCGCACTCACTTGTGGCTTCGATCCTACCGCTG ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA ATTATGACTGGTTCAGCAATATGAATGTGCTGACCTTCCTGCGCGATAT TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCCC TACAACCTGCTGCAGGGTTATACGTATGCCTGTCTGAACAAACA
SEQ ID NO.: 11	pOMe-9 (active site) Synthetase polynucleotide	CGGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA CTGGCGCAAGGCCCGATCGCACTCACTTGTGGCTTCGATCCTACCGCTG ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC TACAACCTGCTGCAGGGTTATTCGTATGCCTGCGAACAAACA
SEQ ID NO.: 12	pOMe-10 (active site) Synthetase polynucleotide	CGGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA CTGGCGCAAGGCCCGATCGCACTCACTTGTGGCTTCGATCCTACCGCTG ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC CCGTTCCTCGATTTCGACTGTGGAGAAAAACTCTGCTATCGCGGCCAATA

-		ATTATGACTGGTTCAGCAATATGAATGTGCTGACCTTCCTGCGCGATAT TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC TACAACCTGCTGCAGGGTTATACGTATGCCTGTCTGAACAAACA
SEQ ID	pOMe-11	CGGGGGCTGGTACCcCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
NO.: 13		CTGGCGCAAGGCCCGATCGCACTCCTTTGTGGCTTCGATCCTACCGCTG
NO.: 13	(active site)	
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
ł		TACAACCTGCTGCAGGGTTATTCTATTGCCTGTTCGAACAAACA
		GTGTG
SEQ ID	pOMe-12	CGGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
NO.: 14	(active site)	CTGGCGCAAGGCCCGATCGCACTCGTGTGTGGCTTCGATCCTACCGCTG
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
	polynaciconac	CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
1		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATAGTATTGCCTGTTTGAACAAACA
		GTGTG
SEQ ID	pOMe-13	CGGGGGCTGGTACCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGA
NO.: 15	(active site)	CTGGCGCAAGGCCCGATCGCACTCGTGTGTGGCTTCGATCCTACCGCTG
11015	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
	polynacieotide	CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATAGTATTGCCTGTTTGAACAAACA
		GTGTG
SEQ ID	pOMe-14	CGGGGGCTGGTAGCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGA
NO.: 16	(active site)	CTGGCGCAAGGCCCGATCGCACTCTGGTGTGGCTTCGATCCTACCGCTG
11010	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
1		CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
	polynucleotide	
1		CTGATTGGCGACCCGAGCTTCAAGGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATT
		GTTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATATGCGTGCCTGTGAGAACAAACA
		GTGTG
SEQ ID	p-acetylPhe-1	CGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
		CTGGCGCAAGGCCCGATCGCACTCATTTGTGGCTTCGATCCTACCGCTG
NO.: 17	(active site)	
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		<del> </del>

SEQ ID NO.: 18	pBenzophenon -1 (active site) Synthetase polynucleotide	CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGGTCAGGGGATTTCGTTCACTGAGTTTTCC TACAACCTGCTGCAGGGTTATGGTATGG
SEQ ID	pBenzophenon	GCGTTAGCAGAGCGACTGGCGCAAGGCCCGATCGCACTCGGGTGTGGC
NO.: 19	e-2 (active	TTCGATCCTACCGCTGACAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCA
	site) Synthetase	ATGCCTGAAACGCTTCCAGCAGGCGGGCCACAAGCCGGTTGCGCTGGT AGGCGGCGCGACGGGTCTGATTGGCGACCCGAGCTTCAAAGCTGCCGA
	polynucleotide	GCGTAAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTGG
	polyndereonde	CCGTAAGCAGGTTGCCCCGTTCCTCGATTTCGACTGTGGAGAAAACTCT
		GCTATCGCGGCCAATAATTATGACTGGTTCGGCAATATGAATGTGCTGA
		CCTTCCTGCGCGATATTGGCAAACACTTCTCCGTTAACCAGATGATCAA
		CAAAGAAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTC
		GTTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATGGTTATGCCTGTA
		TGAACAAACAGTACGGTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTG GGGTAACATCACTTCTGGTATCGACCTGACCCGTCGTCTGCATCAGAAT
		CAGGTG
SEQ ID	pAzidoPhe-1	GGGCTGGTAGCCCAGGTGACGGACGNAGAAGCGTTAGCAGAGCGACTG
NO.: 20	(active site)	GCGCAAGGCCCGATCGCACTCCTTTGTGGCTTCGATCCTACCGCTGACA
	Synthetase	GCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAG
	polynucleotide	CAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGACGGGTCTG
		ATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAA
		GAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCG
		TTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATAATT ATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATATTGG
		CAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAAGCA
		GCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCCTAC
		AACCTGCTGCAGGGTTATTCTATGGCCTGTGCGAACAAACA
		TGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTCTGG
		TATCGACCTGACCCGTCGTCTGCATCANAATCANGTG
SEQ ID	pAzidoPhe-2	TTAGCAGAGCGACTGGCGCAAGGCCCGATCGCACTCGTTTGTGGCTTCG
NO.: 21	(active site) Synthetase	ATCCTACCGCTGACAGCTTGCATTTGGGGCCATCTTGTTCCATTGTTATGC CTGAAACGCTTCCAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGC
	polynucleotide	GGCGCGACGGGTCTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGT
	porymuciconde	AAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGT
		AAGCAGGTTGCCCCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTA
		TCGCGGCCAATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTT
		CCTGCGCGATATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAA
		GAAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTC
		ACTGAGTTTTCCTACAACCTGCTGCAGGGTTATTCTGCGGCCTGTGCGA ACAAACAGTACGGTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGG
		GTAACATCACTTCTGGTATCGACCTGACCCGTCGTCTGCATCAGAATCA
L	<u> </u>	GIAACATCACTICIOGIATCOACCIGACCCGICGICTCAGAATCA

SEQ ID   No.: 22   Synthetase   polynucleotide   Security   Synthetase   Synthetase   Synthetase   Synthetase   polynucleotide   Security   Synthetase   Synthetase   Synthetase   Synthetase   Synthetase   Security   Securit			GGTG
Sec   D   PazidoPhe-5   Control Cont	SEO ID	pAzidoPhe-3	
Synthetase polynucleotide GGCGTGGTGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG			
polynucleotide  GCGTGGTAGGCGGCGCGACGAGAGAACTGTTCAGGGTGGTGGGGGGCGCACAAAAATCCGTAAGCCGGTAACCCGAAGAAAATCTTCTAGGAGTGGGTGG	11022		
GCTGCGAGCGTAAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTGACACATATCGTAACCGATTCGCACTGTTCGACTTGGAGAAAACTCTGCTATTCGACTGTGGAGAAAAATTATGACTGGTTCGGCAATATGACTGGTTCGGCAATATGACTGGTTCGGCAATATGACTGGTTCACCAGGGTTAACCAGAATATATAT			
GACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATTTCGACTGTGGAA   AAACTCTGCTATCGCGCCCATTAATTATGACTGGTTGGCAATATGAA   AGACTCTCCTGCTATCGCGGCCAATATTATGACTGGTTGGGCAATATGAA   GGGATTTCGTTCACCGGCGATATTGGCAAACACTCTCCCGTTAACCAG   GGGATTTCGTTCACCAGGTTTACCAGCGTTCAACCGTGAAGACTCTCTGCGCGAGTTATTCGGC   TGCCTGTGCGAACAAAACAGTAACGGNGNGGNGCTGCAAATTGGNGGTTC   TGACCAGGGGGTAAACACACTCCTTCTGGTATTCGACCTTGCTCTGCAGCGTTATTCGGC   TGCCTGAGACAACAACACTACATCTCTGGTATTCGACCTTGCTCTGCACCGTTATCTGGCT   TGACCAGGGGGGTAAACACACTACTCTTGTGGTATTCGACCTTGCTCTGCACCGTTCTCTCAAAATCAGGTG   SEQ ID   PaidoPhe-5     GCGTTAGCAGCGGGCACAAGCCCGAACGCACTCGTTTGTGCACACTGCTTCTCGCGAAGCGTTAACCAGTTTCGCGAACACTTCCGGTTAACCAGTTTCGCGCAAAATCAGGTTGCCTGAAACCGTTCACACGTTTCGCGCAAAATCAGTGCCGAACACACTCACT		porynacieonae	
AAAACTCTGCTATCGCGGCCAATATGCATGGGTTCGGCAATTAGAG TGTGCTGACCTTCCTGGCGCATATTGGCAAACACTTCTCGGTTAACCAG ATGATCAACAAANAAGCGGTTAGCAGCACTTCTCGGTTAACCAG GGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGCAAGGTTATCGGC TGCCTGTGCGAACAAACAGTTCTCTACACCTGCAAGGTTATTCGGC TGCCTGTGCGAACAACACAGTTCCTACCTGCTAACCCGTCGAACTCGT TGACCAGGGGGGTAACATCACTCTTGTGTATCGACCTGCACCCTTCGTCT TGACCAGGGGGGTAACATCACTCTTGTGTATCGACCTGCACCCTTCTTTGTGGC ATCAAAATCAAGTG  SEQ ID PAzidoPhe-4 (active site) Synthetase polynucleotide GCGGTCGGCGACAGCTTCGACAGCTTGCACTCGTTTGTGGC CTATCGCGGCAACGGTTCACACCGTTCATTGTGCGCTGTAACCAGCTGACCCGACCGA			
TGCTGACACAANAACGGTTAAACACCTCCACGTGAACACCACACACACACAC	1		
ATGATCAACAAANAAGCGGTTAAGCAGGGTTCACACCGTGAACATCAG GGGATTTCGTTCACTCACACCTGCTGCAGGGTTATTCGGC TGCCTGTGCGAACAAACAGTACGGNGNGGNGCTGCAAATTGGNGGTTC TGACCAGGGGGGTAACATCACTTCTGGTATCCACCCTGACCCGTGTCTC CATCAAAATCAGGTG  SEQ ID No.: 23  SEQ id clive site Synthetase polynucleotide  PazidoPhe-4 (active site) Synthetase polynucleotide  ACTACAACAGCGACTGGCGCCAAGGCCCGATCGCACTCGTTTGTGGCT CGTAGCCAGCGCGCACAGGCGGGCCACAAGCCGGTTGCACTGTAGCACGTGAACCGTCAACCGTGAAACCGTTCAAACGGCGACCGAACCCGGTTAGCCGCAAACCCGAAACCCGAAACCCGGTAAGCCGGTAAGCCGGTAAGCCAGCTCAAAACACCTCCACCGTGAAACCCCGAACCCGGCAACACCCGGTTGCACCGTAAGCCGGCGACCGAAAACCCTTCCAGCAAACCCGAAACACCTTCCAGCAAACCCGAAAACCCGAAAACCCGAAAACCCGAAAAACCATTCCCGCGAAAACACCTTCCAGCAAACCCGAAAACCCGAAAAACCCTTCCCGAAAACCCTTCCAGCAAAACCCTTCCAGCAAAACCCTTCCAGCAAAACCCTTCCAGCAAAATCCCTTCCGCGCAAATTGGAAAAACACTTCCCGCTTAACCAGAAAAACACGTTTCCAGCAGAAAACCGTTCCAACCGTGAAAATGTGCCGACCTGTTTCCAGCAGGAAAAACACTTCCACCGTGAAAATCGAGTTCAAACCAAAACAAAC			
SEQ ID pAzidoPhe-4 (active site) Synthetase polynucleotide Physiolates Physiolate			
SEQ ID NO.: 24 PAZIdoPhe-5 (active site) Synubclase polynucleotide			
SEQ ID No.: 24 (active site) synthetase polynucleotide polynucleot			
SEQ ID   PAzidoPhe-4 (active site)   Synthetase   polynucleotide   GGTTAGCAGACCGTTCCACCGTGACAGCTTGCACTTGTTGTGCT   TCGATCCTACCGGCGGGCCACAAGCCGAGCTGGACAAATCCGTGACAAATCCGTACAAACCCGAGCGGGCCACAAGCCGGTGGACAAATCCGTACAAACCCGAACAAATCCGTTCCACAAAACCCGAACAAAACCCGAACAAATCCGTTCCTGCGGCGAACAACCGAACAAATCCGTTCCTGCGGCGAACAACCGAACAACCCGAACAACCCGAACAACCCGAACAAC			
SEQ ID No.: 23   PacidoPhe-4 (active site) Synthetase polynucleotide   GCGTTAGCAGCACTGGCCAAAGCCGGCACAAGCCGGTTGCACTGTTGTGCCAGCAGCTTGCACTTGTTCCATTTGTTCCATTTGTTCCAGCAGCTTGCAGCAGCTTGCAAAGCTTGCAAAGCTTGCAAAGCTGCCGAAAGCCGGTTGCAAAGCTGCCGAAAGCCGGTTGCAAAGCTGCCGAGCCGAAACCGGTTGCAAAGCTGCCAAAACCGCGAAAACCGTTCAAAACCTGCCGAAAACCGCGTAAAGCCTGCCAAAACCGCGAAAACCGTTCAAAACCTGCAAAACCTGCAAAACCACCGAAAAAAACCAGTTCAACCGGAAAAACCACAAAACAAAC			TGACCAGGGGGTAACATCACTTCTGGTATCGACCTGACCCGTCGTCTG
NO.: 23   Synthetase polynucleotide   Synthetase polynucle			CATCAAAATCAGGTG
NO.: 23   Synthetase polynucleotide   Synthetase polynucleotide   Synthetase polynucleotide   GCGGGGGGGGGGGGGGGGGCACAAGCCGACAGGGGGGGG	SEQ ID	pAzidoPhe-4	GCGTTAGCAGAGCGACTGGCGCAAGGCCCGATCGCACTCGTTTGTGGCT
Synthetase polynucleotide    FGCTGAAAACGCTTCCAGCAGGGGGGCCACAAGCCGGTTGCGTGATA GGCGGCGCGACGGGTCTGATTGGCGACCCGAGCTTCAAAGCTGCCAG CCTAAGCTGAACACCGAAGAAAACTGTTCAGGAGTTGGACAAAAACCCGTAAGCAGGTTGCCCCGTTCCTCGATTTCGACATGTGAGAAAAACTCCTTCTCTGCGCGAAAAATTTAGACTGGTTCGACATTGAAACACGAAAAACCTTCCTGCGCATAAGCTAGATTAGACAGGATTTCCAACCGTGAAGAAAACTCTCTACTGAGTTTTCAACAAACA			TCGATCCTACCGCTGACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTG
Polynucleotide   GGCGGCGCGACGGGTTCGATTGGGACCCCGAGCTTCAAAGCTGCCGAG   CGTAAGCACCGAAGAAACTGTTCAGGAGTGGGTGGACAAAATC   CGTAAGCAGCTTGACACCGAAGAAACTGTTCAGGAGTGGGTGG			
CGTAAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTGG			
SEQ ID NO: 25 NO: 26 NO: 26 NO: 26 NO: 27 NO: 26 NO: 26 NO: 27 NO: 28 NO: 26 NO: 25 NO: 26 NO: 25 NO: 26 NO: 25 NO: 26 NO: 27 NO: 26 NO		porymatrical	
CTATCGCGGCAATAATTATGACTGCTTCGCGAATTATGAATGCTGAC CTTCCTGCGCGATATTGCAAACACTTCTCCGTTAACCAGATGATCAAC AAAGAAGCGGTTAAACCAGCTCTCAACCGTGAAGATCAACAC AAAGAACCGGTTAAACCAGCTCTCAACCGTGAAGATCACAGGGGATTTCG TTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAGTGCGGCCTGTGT TAACAAACAGTACGGTTGTGTCTGCACAATTGGTGGTTCTGACCAGTGG GGTAACATCACTTCTGGTATCGACCCGTCGCTGCATCGACACT ANGTG  SEQ ID NO.: 24 Synthetase polynucleotide  GCCTGGTAGCGGCGCAAGGCCGACGGCCACAGCCGGTT GCCTGAGACGCTTCAAACCGTGAAACCGTTCAACACTC GCCTGGAGCGTAGCCGACCGTTCCACCACAGCCGGTT GCCCGAGCGTAGCCGACCGGTCGCACACACCGGTT GCCCGAGCGTAAGCTGAAACCGTTCAACACTGCTCAAACCTTCAACACTGCTCAAA TGTGCTGACCTTCCTGCGCGCAATGATATGAACACGGTTCAACACTGCTCAACCGTTCACCAG ATCAGAATCAGACAAAACAGAAACACGTTTCCATCCGTTAACCAG ATCAGAATCAGGTG  SEQ ID NO.: 25 SQ ID No.: 26 QCCCGAGCGCAACAACACCGAGCCGACCCGACCTCACCGTCGCACCCTCGCACCTTCCCGCACCGACTTCCCGCACCCGACCTCCCGACCTCCCGACCTCCCCGACCTCCCCACCCTCCCCCCCC			
CTTCCTGCGCGATATTGGCAAACACTTCTCCGTTAACCAGATGATCAAC AAAGAAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAACCACAAAGAAGCAGTTTTCCTTACAACCTGCTGCAAGCGTTATAGTGCGGCCTGTGT TAACAAAACAGTACGGTGTGGTGCTGCAAAATTGGTGGTTCTGACCAGTGG GGTAACATCACTTCTTGGTATCGACCTGCACGGTTATAGTGCGGCCTGTGT AACAAACAGTACCGTTTGGATCGACCTGCACTCGCATCGATCAACACTACATTCTGGTATCGACCTGCACCGTCGCTCGC			
AAAGAAGCGGTTAAGCAGCGTCCAACCGTGAAGATCAGGGGATTCG TTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAGTGCGGCCTGTGT TAACAAACAGTACGGTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGG GGTAACATCACTTCTGGTATCGACCTGACCCGTCGTCTGCATCAGAATC ANGTG  SEQ ID DAzidoPhe-5 (active site) Synthetase polynucleotide  GCGCTGGTAGCGCGCAAGGCCCGATCGCACTC GCGCGAGGCGAACGCGCCAAGCCCGACCGGTT TCCATTGTTATGCCTGAAACGCTTCACCGCTGACAGCCCGACCCGACCGGTT GCACAAACCGTTAGCAGCGCGCAAGCCGGTTCAAAA GCTGCCGAGCGCAAACCCGAAGAAACCGTTCAACACAGAAAATCGTTAACAG AAAACCCTTACCAGGCGCAATGTTCCAGCAGCCGGCCAAAACCGGTT GACAAAAATCCGTAAACAAAAACACTATCCGCGAAAACACTTCTCCGTTAACCAG ATGATCAACAAAAGAAACACTTCCACGCGCAAATTGGAA ATGATCAACAAAAGAAACACTTCTCGACTTCCGCGAAATTGAA TGTGCTGAACCAAGAAAACACTTCTCGCTTCACCGTGAAGAATATTT TGCCTGTGTGAACAAAAAAACACTTCTCGCTTCAACCGTGAAGATCAG GGGATTTCGTTCACTGAGCTGTTCTCAACCTGCTGCAACGTTCTC GACCAGTGGGTAACAACAACACTTCCGCTGCAACGTTCCACCG TTCCAGCAAGCAGCTAACACTTCTCGCTAACCGTGACCCGTCCTCTC ATCAGAATCAGGTG  SEQ ID pAzidoPhe-6 (active site) Synthetase polynucleotide  CGCTGGTGAACAAACACTTCTCGGTATCGACCCGTCCACCGTCCACCGTCCTCACCG CTGACAACACACACACCAAGCCCGTTCCAACCGTTCACCGCACAC ATATTATGACCTGACC			
TTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAGTGCGGCCTGTGT TAACAAACAGTACGGTGTGTGCTGCAAATTGGTGGTTCTGACCAGTGG GGTAACATCACTTCTGGTATCGACCTGACCCGTCTGCATCAGAATC ANGTG  SEQ ID NO.: 24  ACITIC Synthetase polynucleotide  GGCTGGTAGGCGGCGAAGGCCCGATCGACCCGTCTGCATCAAACCCGTTGTTATTGCGGGCATCTAAAACCCGTTAACAACACAAAAATCCGTAACACCAGGAAAACACTTCTCGGTTAACAACAGGAAAAATCCGTAACACCAGGAGAAACACTGTTCAGCAAAACAGAAAATCAGTTCAGCAAAACACAGAAAAATCAGTTCAACAAAAAAACAGTACCGAAGAAACACATCACCGCGAAGAACACCTTCAACCGCGAAGAAACCGTTCAACACCGAGGAAAACCAGTTAAATTT TGCCTGGACAAACAAGAAACAGTTCCACCAAAAATCAGTACACAAAAAAAA			
SEQ ID pAzidoPhe-5 (active site) Synthetase polynucleotide  SEQ ID NO: 24  SEQ ID PAzidoPhe-5 (active site) Synthetase polynucleotide  Polynucleotide  SEQ ID PAzidoPhe-5 (active site) Synthetase polynucleotide  Polynucleot			
SEQ ID NO: 24 PAzidoPhe-5 (active site) Synthetase polynucleotide (active site) Synthetase			
SEQ ID No.: 24 (active site) Synthetase polynucleotide			
SEQ ID NO.: 24 Synthetase polynucleotide PAzidoPhe-5 (active site) Synthetase polynucleotide Pol			1
NO.: 24   Synthetase polynucleotide   Synthetase polynucleotide   Synthetase polynucleotide   Synthetase polynucleotide   GCGCTGGTAGGCGGCGACGGGTTCAACAGCAGGCGGCCACAAGCCGGGTT   GCGCTGGTAGGCGGCGACGGGTTCAACACACCGAAGACCCGAAGCTGCAAA   GCTGCCGAGCGTAAGCTGAACACCGAAGAAACCTGTCAGAAAAATCCGTAAGCAGGTTGCCCGTTCCTCGATTTCGACTGTGGAG   AAAACTCTGCTATCCGCGCCAATGATTATGACTGGCAATATGAA   TGTGCTGACCTTCCTGGCCGATATTGGCAAACACTTCTCCGTTAACCAG   ATGATCAACAAAGAAGCGGTTAAGCAGCGTTCAACCAG   ATGATCAACAAAGAAGCGGTTAAGCAGCGTTCAACCAG   ATGATCAACAAAGAAGCGGTTAAACCAGCGGGATTTCTTCACTACAACCTGCACGTGAAGATCAGGTGGAACAAACA			
Synthetase polynucleotide    Synthetase polynucleotide   CCATTGTTATGCCTGAAACGCTTCCAGCAGGCGGGCCCAGAGCTCAAAGCCGGTT			
pólynucleotide GCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGACCCGAGCTTCAAA GCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTTCAGAGTGGAGGTG GACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATTTCAGCAGTGGAG AAAATCCGTAACACCGAAGAAACTGTTCAGCAGTGAGAGAAAATCCGTAACACGCAATTATTGACTGGTAACAACACAAAAAACACTTCTCCGTTAACCAG ATGATCAACAAAAAAAAAA	NO.: 24		
GCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTG GACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATTTCAGCTGGAG AAAACTCTGCTATCGCGGCCAATGATTATGACTGGTTCGGCAATATGAA TGTGCTGACCTTCCTGCGCGATATTGGCAACCGTTCACCAG ATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAG GGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAATTT TGCCTGTGTGAACAACAACACTTCTCGCAGGGTTATAATTT TGCCTGTGTGAACAACACTTCTGGTATCGACCGTCGCAGGGTTATAATTT TGCCTGTGTGAACAACACTTCTGGTATCGACCTGCACCGTCGTCTGC ATCAGAATCAGGTG  SEQ ID NO:: 25 Synthetase polynucleotide CGACTGGCGCAAGGCCCGATCGCACTCACGTGTGGCTTCGAAACGC TTCCAGCAGCGGGCCACAAGCCCGTTGGCTTCGATCCTACCG CTGACAGCTTGATTTGGGGGATCTTCATTCATGCTGAAACGC TTCCAGCAGCGGGCCACAAGCCGGTTGCACAAAATCCGTAAGCAGGTT GCCCCGTTCCTCGATTTCGACTGTGGAAAAAATCCGTAAGCAGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGAAAAATCCGTAAGCAGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGAGAAAAATCCGTAACAAAAACAGTA CGGTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGTTAACAAAACAGTA CGGTGTGGTGCCAAATTGGTGGTTCTGACCAGTGGGTAACACACAAGTA CGGTGTGGTGCCAAATTGGTGGTTCTGACCAGTGGGTTAACAAAACAGTA CGGTTGGTGCCAAAATTGGTGGTTCTGACCAGTGGGTAACACACAC		Synthetase	
SEQ ID NO.: 25 PAZidoPhe-6 (active site) Synthetase polynucleotide Fig. 1 AAATTATGACGGGCAAATGATCAGGGGCAAATGATCAGCAGGGGGGGG		polynucleotide	
AAAACTCTGCTATCGCGGCCAATGATTATGACTGGTTCGCAATATGAA TGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTCTCCGTTAACCAG ATGATCAACAAAGAAGCGGTTTAAGCAGCTCTCACCGTGAAGATCAG GGGATTTCGTTCACTGAGTTTACCTGCAGGGTTATATTT TGCCTGTGTGAACAAACAGTACGGTGTGGTGCTGCAAATTGGTGGTTCT GACCAGTGGGGTAACATCACTTCTGGTATCGACCTGCTGCCAAATTGGTGGTTCT GACCAGTGGGGTAACATCACTTCTGGTATCGACCTGACCCGTCTCC ATCAGAATCAGGTG  SEQ ID NO.: 25  Spynthetase polynucleotide CGACTGCCGAACCCGATCGCACTCACGTGTGGCTTCGATACCGC ACCAGCGCGCACGGGCCACAAGCCGGTTGCCATTGTTATGCCTGAAACGC TTCCAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGAAG ACCGAAGAAACTGTTCAGGAGTGGGTAAACTCTGAACCGCAATATTATGACTGGTTCCGATTCGGCAAAAACTGTTAACAAAACAGTT GCCCCGTTCCTCGATTTCGGCAATATGAATGACTGCTAACCGCCAATATGAATCACTTCCTGCTGAAACCGCTTCCAAACCGTTAACCAGATGATCAACAAAAAAAA	Į.		GCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTG
TGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTCTCCGTTAACCAG ATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAG GGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAATTT TGCCTGTGTAACAAACAGTACGGTGGTGCTGCAAATTGGTGGTTCT GACCAGTGGGGTAACATCACTTCTGGTATCGACCTGACCCGTCGTCG ATCAGAATCAGGTG  SEQ ID NO.: 25  Spinthetase polynucleotide  PAzidoPhe-6 (active site) Synthetase polynucleotide  ACCAGCGGGCGCAAGGCCCGATCGCACTCACGTGTGGCTTCGATCCTACCG CTGACAGCTTGCATTTTGGGGCATCTTGTTATGCCTGAAACGC TTCCAGCAGGCGGGCCACAAGCCGGTTCAAAGCTGCAGCGGACG ACCGAAGAAACTGTTCAGGAGTGGGTGAGCACGAAC ACCGAAGAAACTGTTCAGGAGTGGGTGAACACTTCCTGCGCGA ATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCA ATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCA TATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAAAAACAGTT AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAAGGTTATTCGGCTGCCTGTCTTAACAAACA	ŀ		
ATGATCAACAAGAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAG GGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAATTT TGCCTGTGTGAACAACAGTACGTTGTGTGTGCTGCAAATTGGTGGTTCT GACCAGTGGGGTAACATCACTTCTGGTATCGACCCGTCGCAAATTGGTGGTTCT ATCAGAATCAGGTG  SEQ ID NO.: 25 Synthetase polynucleotide Synthetase polynucleotide ACGACAGCGGCCCGACCCACCCACTCACGTGTGAGCCCGACC GGTCTGATTGCACTTTGTTCCATTGTTATGCCTGAAACCC TTCCAGCAGCCGAGCCCGAGCTTCCATTGTTATGCCTGAAACCC GGTCTGATTGCACCCGAGCTTCAAACCTGCCGACCGACGCGACG ACCGAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGCA ACCGAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAACCAGAGCAAAACACTTCCTCCGCCA ATAATTATGACTGGTTCGGCAATATGAATGATCGTCACCTGCCGAG TATTGGCAAACACTTCTCCGTTAACCAGATGATCACAAAAAAAA			AAAACTCTGCTATCGCGGCCAATGATTATGACTGGTTCGGCAATATGAA
SEQ ID NO.: 25 PAZIGOPhe-6 (active site) Synthetase polynucleotide ATATTATGCCGAACCGTCGCAACCGTCGCAACCGTCGCAACCGTCGCAACCGTCGCAACCGTCGCACCGTCGCAACCGTCGCACCGTCGCAACCGACCG			TGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTCTCCGTTAACCAG
SEQ ID NO.: 25 SEQ ID NO.: 25 SEQ ID NO.: 25 SEQ ID NO.: 25 SEQ ID NO.: 26 SEQ ID			ATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAACCGTGAAGATCAG
SEQ ID NO.: 25 PAZIDOPhe-6 (active site) Synthetase polynucleotide ATAATTATGACCGAACCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCAACCCGAAGAA			GGGATTTCGTTCACTGAGTTTTCCTACAACCTGCTGCAGGGTTATAATTT
SEQ ID NO.: 25 PAZIDOPhe-6 (active site) Synthetase polynucleotide ATAATTATGACCGAACCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCCCGAACCAACCCGAAGAA			TGCCTGTGTGAACAAACAGTACGGTGTGGTGCTGCAAATTGGTGGTTCT
SEQ ID NO.: 25 SEQ ID Synthetase polynucleotide Polynucleotide ACCGAAGAACCCGACCGACCGACGACGCGACGAGGCCGACGA			
SEQ ID NO.: 25  Synthetase polynucleotide			
NO.: 25 (active site) Synthetase polynucleotide  FOR CAGAGAGAGAGAGCAGGAGCATCTTGTTCCATTGTTATGCCTGAAACGC  Synthetase polynucleotide  FOR CAGAGAGAGAGCAGGCACAAGCCGGTTGCGCTGGTAGGCGGCGACG GGTCTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCAGACAAAAAAAA	SEO ID	nAzidoPhe-6	
Synthetase polynucleotide GGTCTGATGGCGGGCGACGGGCGACGGGTTGCGCTGGTAGGCGGCGCGACGGGTTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTTCAGGAGTGGGTGG		_	CTGACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGC
polynucleotide GGTCTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAAC ACCGAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTT GCCCCGTTCCTCGATTTCGACTGTGGAGAAAAACTCTGCTATCGCGGCCA ATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGA TATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA	110 23		
ACCGAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTT GCCCCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCA ATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGA TATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA			
GCCCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCA ATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGA TATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA		polynacieonae	
ATAATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGA TATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA			
TATTGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTT AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA			
AAGCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTT CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA	ļ		
CCTACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA			
CGGTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACT TCTGGTATCGACCTGACCGTCGTCTGCATCAGAATCAGGTG  SEQ ID pPR-EcRS-1 (propargyloxy phenylalanine synthetase) (active site) Synthetase Synthetase GAAGAAACTGTTCAGGAGCCCGATCGCACTGGTTGCTTCAAAGCTGCACTGAAACCCCGTTGATCCTACAGGTTGCATTTGTTCAAAGCTGCAAACCCCGGTTGCATCTAGAACCCCGTTGATTGTAGCAGAACCCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCCGAAGAAAATCCGTAAGCAGGTTGCCC			
SEQ ID pPR-EcRS-1 CGGGGGCTGGTANCCCAGGTGACGGAGGAAGCGTTAGCAGAGCGA (propargyloxy phenylalanine synthetase) (active site) Synthetase GAAGAAACTGTTCAGGAGCTGACCGAGGTTAGCAGAGCGTTAGCAGAGCCTTCAAAGCTTCAGAGCGGACGAGCGGAAGCCTTCAAAGCTTCAAAGCTTCCATTGTTATGCCTGAAACCCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACCCGAGAGAAAACTGTTCAGGAGTGGACAAAAATCCGTAAGCAGGTTGCC			
SEQ ID pPR-EcRS-1 CGGGGGCTGGTANCCCAGGTGACGAGGAAGCGTTAGCAGAGCGA (propargyloxy phenylalanine synthetase) (active site) Synthetase Synthetase GAAGAAACTGTTCAGGAGTGGCCGACGAGGCGACGAGGCGACGAGCCGACCGA			
NO.: 26 (propargyloxy phenylalanine synthetase) (active site) Synthetase Synthetase (AGAGAAAACTGTTCAGGACCGACTCGAGTTGCATTGTTATGCCTGAAACGCTTCAAAGCTGCACTGCACTTGTTATGCCTGAAACGCTTCAAAGCTGCACTGTAGGCGGCGACGGGTTGCAAAGCTGCCGAGCTTCAAAGCTGCCGAGCTTAAGCAGCTGAACACCCAGAAAACTGTTCAGGAGTGGGTGG			
phenylalanine synthetase) (active site) Synthetase  ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC			
synthetase) CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGACGGGT (active site) CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC	NO.: 26		
(active site) CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC Synthetase GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC	-		
Synthetase GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC	i		
Synthetase GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC	1	(active site)	
	1		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
, , , , , , , , , , , , , , , , , , , ,	1	polynucleotide	CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA

		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATTCTATGGCCTGTTTGAACAAACA
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGANCCGTCGTCTGCATCAGAATCAGGTG
SEQ ID	pPR-EcRS -2	CGGGGCTGGTAGCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGA
NO.: 27	(active site)	CTGGCGCAAGGCCCGATCGCACTCACGTGTGGCTTCGATCCTACCGCTG
1102.	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
	porynaciconac	CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		•
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGGATTTCGTTCACTGAGTTTTCC
		TACAATCTGCTGCAGGGTTATTCGGCTGCCTGTCTTAACAAACA
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGAACCTGANCCGTCGTCTGCATCAAAATCAAGTG
SEQ ID	pPR-EcRS -3	CGGGGCTGGTACCCCAAGTGACGGACGAGGAAACGTTAGCAGAGCGA
NO.: 28	(active site)	CTGGCGCAAGGCCCGATCGCACTCTCTTGTGGCTTCGATCCTACCGCTG
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCAGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATACGATGGCCTGTGTGAACAAACA
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGACCCGTCGTCTGCATCAGAATCAGGTG
SEQ ID	pPR-EcRS -4	CGGGGCTGGTAGCCCAGGTGACGACGAGGAAGCGTTAGCAGAGCGA
NO.: 29	(active site)	CTGGCGCAAGGCCCGATCGCACTCGCGTGCGGCTTCGATCCTACCGCTG
110 25	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGACGGGT
	porynacieonae	CTGATTGGCGACCCGAGCTTCAAGGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATTCTTATGCCTGTCTTAACAAACA
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGACCCGTCGTCTGCATCAGAATCAGGTG
SEQ ID	pPR-EcRS-5	CGGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
NO.: 30	(active site)	CTGGCGCAAGGCCCGATCGCACTCGCGTGTGGCTTCGATCCTACCGCTG
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
	ļ	TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATACGATGGCCTGTTGTAACAAACA
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGACCCGTCGTCTGCATCAGAATCAGGTG
SEQ ID	pPR-EcRS-6	CGGGGGCTGGTACCCCAAGTGACGGACGAGGAAGCGTTAGCAGAGCGA
DEQ ID	PLK-ECKO-0	COOCCIOTACCCAAOTOACOGACOAOOAAOCOTTACCAOAOCOA

NO.: 31	(aatiua aita)	CTGGCGCAAGGCCCGATCGCACTCACGTGTGGCTTCGATCCTACCGCTG
NO.: 31	(active site)	
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCGCTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATACGTTTGCCTGTATGAACAAACA
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGACCCGTCGTCTGCATCAGAATCAGGTG
CEO ID	DD C.DC 7	GTGACGGACGAGGAAGCGTTAGCAGAGCGACTGGCGCAAGGCCCGATC
SEQ ID	pPR-EcRS-7	
NO.:32	(active site)	GCACTCACGTGTGGCTTCGATCCTACCGCTGACAGCTTGCATTTGGGGC
	Synthetase	ATCTTGTTCCATTGTTATGCCTGAAACGCTTCCAGCAGGCGGGCCACAA
	polynucleotide	GCCGGTTGCGCTGGTAGGCGGCGCGACGGGTCTGATTGGCGACCCGAG
		CTTCAAAGCTGCCGAGCGTAAGCTGAACACCGAAGAAACTGTTCAGGA
		GTGGGTGGACAAAATCCGTAAGCAGGTTGCCCCGTTCCTCGATTTCGAC
		TGTGGAGAAAACTCTGCTATCGCGGCCAATAATTATGACTGGTTCGGCA
·	i	ATATGAATGTGCTGACCTTCCTGCGCGATATTGGCAAACACTTCTCCGT
1		TAACCAGATGATCAACAAAGAAGCGGTTAAGCAGCGTCTCAACCGTGA
		AGATCAGGGGATTTCGTTCACTGAGTTTTCCTACAATCTGCTGCAGGGT
		TATTCGGCTGCCTGTCTTAACAAACAGTACGGTGTGGTGCTGCAAATTG
		GTGGTTCTGACCAGTGGGGTAACATCACTTCTGGTATCGACCTGACCCG
	77 7 7 7 7	TCGTCTGCATCAGAATCAGGTG
SEQ ID	pPR-EcRS-8	CGGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
NO.: 33	(active site)	CTGGCGCAAGGCCCGATCGCACTCGTTTGTGGCTTCGATCCTACCGCTG
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
1		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
1		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
		TACAACCTGCTGCAGGGTTATTCGATGGCCTGTACGAACAACAGTACG
		GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGACCCGTCGTCTGCATCAGAATCAGGTG
OEO ID	DD E DC O	CGGGGCTGTANCCCAAGTGACGGACGGGAAGCGTTAGCAGAGCGA
SEQ ID	pPR-EcRS-9	
NO.: 34	(active site)	CTGGCGCAAGGCCCGATCGCACTCAGTTGTGGCTTCGATCCTACCGCTG
	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
1	1	CCGTTCCTCGATCTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
1	1	ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
1		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA
1		GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC
1	1	TACAACCTGCTGCAGGGTTATAGTTTTGCCTGTCTGAACAAACA
	1	GTGTGGTGCTGCAAATTGGTGGTTCTGACCAGTGGGGTAACATCACTTC
		TGGTATCGACCTGACCCGTCGTCTGCATCAGAATCAGGTG
CEO ID	aDD FaDC 10	CGGGGCTGGTAGCCCAGGTGACGGACGAGGAAGCGTTAGCAGAGCGA
SEQ ID	pPR-EcRS-10	
NO.: 35	(active site)	CTGGCGCAAGGCCCGATCGCACTCACGTGTGGCTTCGATCCTACCGCTG
1	Synthetase	ACAGCTTGCATTTGGGGCATCTTGTTCCATTGTTATGCCTGAAACGCTTC
1	polynucleotide	CAGCAGGCGGCCACAAGCCGGTTGCGCTGGTAGGCGGCGCGACGGGT
1		CTGATTGGCGACCCGAGCTTCAAAGCTGCCGAGCGTAAGCTGAACACC
		GAAGAAACTGTTCAGGAGTGGGTGGACAAAATCCGTAAGCAGGTTGCC
		CCGTTCCTCGATTTCGACTGTGGAGAAAACTCTGCTATCGCGGCCAATA
		ATTATGACTGGTTCGGCAATATGAATGTGCTGACCTTCCTGCGCGATAT
	•	

	r	TOO A A CACTTOTOCCTTA ACCAC ATCATCA ACCAA CACAA COCCTTA A
		TGGCAAACACTTCTCCGTTAACCAGATGATCAACAAAGAAGCGGTTAA GCAGCGTCTCAACCGTGAAGATCAGGGGATTTCGTTCACTGAGTTTTCC TACAACCTGCTGCAGGGTTATACGTTTGCCTGTACTAACAAACA
SEQ ID	p-iodoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALVCGFDPTADSLHLGH
NO.: 36	Synthetase Amino acid (aa)	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM INKEAVKQRLNREDQGISFTEFSYNLLQGYSYACLNKQYGVVLQIGGSDQ WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
SEQ ID	p-iodoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALICGFDPTADSLHLGH
NO.: 37	2 Synthetase Amino acid	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACLNKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
SEQ ID	p-iodoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALVCGFDPTADSLHLGH
NO.: 38	3	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
İ	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACANKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
1		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
SEQ ID	OMeTyrRS-1	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALVCGFDPTADSLHLGH
NO.: 39	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACLNKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
	0) ( 5 7 7 7 7	GRFTLLRRGKKNYCLICWK
SEQ ID	OMeTyrRS-2	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH
NO.: 40	Synthetase Amino acid	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYTMACLNKQYGVVLQIGGSDQ
	(44)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	OMeTyrRS-3	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH
NO.: 41	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYTYACLNKQYGVVLQIGGSDQ
	'	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	1	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
	1	AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK

SEQ ID	OMeTyrRS-4	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALLCGFDPTADSLHLGH
NO.: 42	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
110 42	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
		INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACSNKQYGVVLQIGGSDQ
	(aa)	
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	OMeTyrRS-5	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALLCGFDPTADSLHLGH
NO.: 43	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACANKQYGVVLQIGGSDQ
	()	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	OMeTyrRS-6	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH
NO.: 44	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYRMACLNKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMOALVDSELOPSRGOARKTIASNAITINGEKOSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALICGFDPTADSLHLGH
NO.: 45	acetylPheRS-1	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
110 43	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
1	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYGMACANKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	(aa)	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
Į.	,	EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALGCGFDPTADSLHLGH
NO.: 46	benzoylPheRS	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	-1	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Synthetase	INKEAVKQRLNREDQGISFTEFSYNLLQGYGFACANKQYGVVLQIGGSDQ
	Amino acid	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	(aa)	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
	, ,	AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	n-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALGCGFDPTADSLHLGH
NO.: 47	p- benzoylPheRS	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
110.:4/		DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	-2	· ·
	Synthetase	INKEAVKQRLNREDQGISFTEFSYNLLQGYGYACMNKQYGVVLQIGGSDQ
	Amino acid	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	(aa)	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-azidoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALLCGFDPTADSLHLGH
NO.: 48	1	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACANKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
L	1.(27)	The state of the s

,		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
OFO ID	: 1 DI DG	
SEQ ID	p-azidoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALVCGFDPTADSLHLGH
NO.: 49	2	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYSAACANKQYGVVLQIGGSDQ
i	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	(aa)	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-azidoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALLCGFDPTADSLHLGH
NO.: 50	3	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
11050	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYSAACANKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
1		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-azidoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALVCGFDPTADSLHLGH
NO.: 51	p-azidor nerco-	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
NO.: 31	4	
	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYSAACVNKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-azidoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALICGFDPTADSLHLGH
NO.: 52	5	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
110 32		DKIRKQVAPFLDFDCGENSAIAANDYDWFGNMNVLTFLRDIGKHFSVNQM
	Synthetase	
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYNFACVNKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
i		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
1		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	p-azidoPheRS-	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH
NO.: 53	6	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
110 33	_	DAIDAON Y DEL DEDCCENG Y IN Y MINADINEGNI MINI LEI DDIGARTEGNIUM
	Synthetase	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	Amino acid	INKEAVKQRLNREDQGISFTEFSYNLLQGYSAACLNKQYGVVLQIGGSDQ
	(aa)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-1	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALGCGFDPTADSLHLGH
		LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
NO.: 54	Synthetase	PAID NOT Y DELI DEDOCENIO Y Y YND ADDIODE OLIVANTAL MET DEDOCATEGRATON
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACLNKQYGVVLQIGGSDQ
	<i>p</i> -	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
	propargyloxyp	PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
	henylalanine	AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
	synthetase	EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-2	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH

NO.: 55	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
11033	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSAACLNKQYGVVLQIGGSDQ
	(44)	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
SEC ID	-DD E-DC 2	GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-3	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALSCGFDPTADSLHLGH
NO.: 56	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYTMACVNKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-4	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALACGFDPTADSLHLGH
NO.: 57	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSYACLNKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-5	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALACGFDPTADSLHLGH
NO.: 58	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
1	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYTMACCNKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-6	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH
NO.: 59	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYTFACMNKQYGVVLQIGGSDQ
		WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-7	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH
NO.: 60	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSVACLNKQYGVVLQIGGSDQ
	· ·	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM
		EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF
		GRFTLLRRGKKNYCLICWK
SEQ ID	pPR-EcRS-8	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALVCGFDPTADSLHLGH
NO.: 61	Synthetase	LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV
	Amino acid	DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM
	(aa)	INKEAVKQRLNREDQGISFTEFSYNLLQGYSMACTNKQYGVVLQIGGSDQ
	`/	WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS
		PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL
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		AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
SEQ ID NO.: 62	pPR-EcRS-9 Synthetase Amino acid (aa)	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALSCGFDPTADSLHLGH LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM INKEAVKQRLNREDQGISFTEFSYNLLQGYSFACLNKQYGVVLQIGGSDQW GNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTSP YKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVLA EQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEME KGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
SEQ ID NO.: 63	pPR-EcRS-10 Synthetase Amino acid (aa)	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALTCGFDPTADSLHLGH LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM INKEAVKQRLNREDQGISFTEFSYNLLQGYTFACTNKQYGVVLQIGGSDQ WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEEDKNSGKAPRAQYVL AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK
SEQ ID	tRNA/Tyr	
NO.: 64	polynucleotide	AGCTTCCCGATAAGGGAGCAGGCCAGTAAAAAGCATTACCCCGTGGTG GGGTTCCCGAGCGGCCAAAGGGAGCAGACTCTAAATCTGCCGTCATCG ACCTCGAAGGTTCGAATCCTTCCCCCACCACCA
SEQ ID	tRNA/Tyr ·	
NO.: 65		AGCUUCCCGAUAAGGGAGCAGGCCAGUAAAAAGCAUUACCCCGUGGU GGGGUUCCCGAGCGGCCAAAGGGAGCAGACUCUAAAUCUGCCGUCAU CGACCUCGAAGGUUCGAAUCCUUCCCCCACCACCA
SEQ ID NO.: 66	Amber Mutants L3TAG	5'-ATGAAGTAGCTGTCTTCTATCGAACAAGCATGCG-3'
SEQ ID NO.: 67	Amber Mutants I13TAG	5'-CGAACAAGCATGCGATTAGTGCCGACTTAAAAAG-3'
SEQ ID NO.: 68	Amber Mutants T44TAG	5'-CGCTACTCTCCCAAATAGAAAAGGTCTCCGCTG-3'
SEQ ID NO.: 69	Amber Mutants F68TAG	5'-CTGGAACAGCTATAGCTACTGATTTTTCCTCG-3'
SEQ ID NO.: 70	Amber Mutants R110TAG	5'-GCCGTCACAGATTAGTTGGCTTCAGTGGAGACTG-3'
SEQ ID NO.: 71	Amber Mutants V114TAG	5'-GATTGGCTTCATAGGAGACTGATATGCTCTAAC-3'
SEQ ID NO.:72	Amber Mutants T121TAG	5'-GCCTCTATAGTTGAGACAGCATAGAATAATGCG-3'
SEQ ID NO.: 73	Amber Mutants I127TAG	5'-GAGACAGCATAGATAGAGTGCGACATCATCGG-3'
SEQ ID NO.: 74	Amber Mutants S131TAG	5'-GAATAAGTGCGACA <b>TAG</b> TCATCGGAAGAGAGTAGTAG-3'
SEQ ID NO.: 75	Amber Mutants T145TAG	5'-GGTCAAAGACAGTTGTAGGTATCGATTGACTCGGC-3'

SEQ ID NO.: 76	Permissive Site Mutants T44F	5'-CGCTACTCTCCCCAAATTTAAAAGGTCTCCGCTG-3'
SEQ ID NO.: 77	Permissive Site Mutants T44Y	5'-CGCTACTCTCCCCAAATATAAAAGGTCTCCGCTG-3'
SEQ ID NO.: 78	Permissive Site Mutants T44W	5'-CGCTACTCTCCCCAAATGGAAAAGGTCTCCGCTG-3'
SEQ ID NO.:79	Permissive Site Mutants T44D	5'-CGCTACTCTCCCCAAAGATAAAAGGTCTCCGCTG-3'
SEQ ID NO.:80	Permissive Site Mutants T44K	5'-CGCTACTCTCCCCAAAAAAAAAAAGGTCTCCGCTG-3'
SEQ ID NO.: 81	Permissive Site Mutants R110F	5'-GCCGTCACAGATTTTTTGGCTTCAGTGGAGACTG-3'
SEQ ID NO.: 82	Permissive Site Mutants R110Y	5'-GCCGTCACAGATTATTTGGCTTCAGTGGAGACTG-3'
SEQ ID NO.: 83	Permissive Site Mutants R110W	5'-GCCGTCACAGATTGGTTGGCTTCAGTGGAGACTG-3'
SEQ ID NO.: 84	Permissive Site Mutants R110D	5'-GCCGTCACAGATGATTTGGCTTCAGTGGAGACTG-3'
SEQ ID NO.: 85	Permissive Site Mutants R110K	5'-GCCGTCACAGATAAATTGGCTTCAGTGGAGACTG-3'
SEQ ID NO.: 86	p- acetylPheRS-1 Synthetase Amino acid (aa) <sup>a</sup>	MASSNLIKQLQERGLVAQVTDEEALAERLAQGPIALICGFDPTADSLHLGH LVPLLCLKRFQQAGHKPVALVGGATGLIGDPSFKAAERKLNTEETVQEWV DKIRKQVAPFLDFDCGENSAIAANNYDWFGNMNVLTFLRDIGKHFSVNQM INKEAVKQRLNREGQGISFTEFSYNLLQGYGMACANKQYGVVLQIGGSDQ WGNITSGIDLTRRLHQNQVFGLTVPLITKADGTKFGKTEGGAVWLDPKKTS PYKFYQFWINTADADVYRFLKFFTFMSIEEINALEEDKNSGKAPRAQYVL AEQVTRLVHGEEGLQAAKRITECLFSGSLSALSEADFEQLAQDGVPMVEM EKGADLMQALVDSELQPSRGQARKTIASNAITINGEKQSDPEYFFKEEDRLF GRFTLLRRGKKNYCLICWK

<sup>&</sup>lt;sup>a</sup> These clones also contain a Asp165Gly mutation